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Dynamics and Control of Flexible Space Vehicles

P. W. Likins

JET PROPULSION LABORATORY
CALIFORNIA INSTITUTE OF TECHNOLOGY
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Preface

The work described in this revised report was performed under the cognizance of the Guidance and Control Division of the Jet Propulsion Laboratory. The original report, dated February 15, 1969, is obsolete.

The purpose of the report is to document and disseminate information of practical value to the engineer charged with the responsibility for developing an attitude-control system for a flexible space vehicle. It is essentially a treatise on the dynamics of flexible vehicles as viewed by the control systems engineer, and deals with control system design only to the extent of suggesting ways in which vehicle flexibility can be accommodated in both preliminary design and final evaluation of the attitude-control system.

The major portion of the report is concerned with the derivation of equations of motion for the computer simulation of a controlled and nonrigid vehicle, and the development of coordinate transformations that facilitate simulation. Three basic approaches to this simulation are covered: discrete-coordinate methods (Section II), hybrid-coordinate methods (Section III), and vehicle normal-coordinate methods (Section IV).

The discrete-coordinate methods of Section II involve few restrictions or approximations, and in some cases are as general as Newton's laws for the simulation of the dynamic response of a collection of interconnected rigid bodies. The limitations of these methods stem primarily from the difficulty of creating the required mathematical model of a real vehicle without exceeding the practical limits imposed on computation by considerations of budget, schedule, and computer capacity.

The hybrid-coordinate methods of Section III receive the greatest emphasis in this report, both because they appear to be most useful and because they are the least familiar. These methods may be applied only when some portions of the vehicle (flexible appendages) undergo deformations that may reasonably be assumed to remain "small," thereby permitting the transformation to modal coordinates for vehicle appendages. The key feature of this approach, as opposed to the discrete-coordinate method, is the possibility of truncating the matrix of modal coordinates.

The vehicle normal-coordinate methods of Section IV involve transformations of all the kinematic coordinates of the simulation, and not merely the appendage deformation coordinates. These methods are accordingly more limited, and even when applicable, they may require more complex coordinate transformations than the hybrid-coordinate methods would involve. In the simplest cases, however, the vehicle normal-coordinate methods probably afford the most efficient simulation, since they then permit the most severe coordinate truncation.

In addition to the material in this report for use in the simulation of a controlled flexible vehicle, there is material of value in preliminary control system design. Because of the modal coordinates employed in both the hybrid-coordinate methods and the vehicle normal-coordinate methods, the corresponding equations of motion permit simulations of varying dimension and complexity, depending on the degree of truncation. As shown in Section V, one can truncate so severely as to represent a very complex structural appendage by a single modal coordinate, and then manually calculate dynamic transfer functions for use in preliminary control system design.
Acknowledgment

A work of this size inevitably reflects the cooperative effort of many people. Among those who actively supported the initiation of this undertaking were Mr. Boris Dobrotin and Mr. H. Karl Bouvier of the Spacecraft Control Section, and Mr. Ben Wada and Mr. Robert Oliver of the Applied Mechanics Section. The work was conducted under the cognizance of Mr. Bouvier.

The original work described in this report (the hybrid-coordinate method of dynamic simulation) was developed from a suggestion offered by Mr. Paul McHorney of Hughes Aircraft Company in the context of a particular problem. Mr. Austin Gale of Hughes Aircraft Company developed a computer simulation program that was useful in the early stages of development of this method.

The technical contributions of JPL personnel to this report are gratefully acknowledged. Of particular value were the critical evaluations of Dr. Marc Trubert, Mr. John Garba, and Mr. Robert Bamford, and the theoretical and computational contributions of Mr. Gerald Fleischer, Mr. Roy Mankovitz, and Mr. Ed Dorroh.

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Abstract

The purpose of this report is twofold: (1) to survey the established analytic procedures for the simulation of controlled flexible space vehicles, and (2) to develop in detail methods that employ a combination of discrete and distributed ("modal") coordinates, i.e., the hybrid-coordinate methods.

Analytic procedures are described in three categories: (1) discrete-coordinate methods, (2) hybrid-coordinate methods, and (3) vehicle normal-coordinate methods. Each of these approaches is described and analyzed for its advantages and disadvantages, and each is found to have an area of applicability.

The hybrid-coordinate method combines the efficiency of the vehicle normal-coordinate method with the versatility of the discrete-coordinate method, and appears to have the widest range of practical application.

The results in this report have practical utility in two areas: (1) complex digital computer simulation of flexible space vehicles of arbitrary configuration subject to realistic control laws, and (2) preliminary control system design based on transfer functions for linearized models of dynamics and control laws.
Dynamics and Control of Flexible Space Vehicles*

I. Introduction

A. Motivation

The development of an attitude-control system necessarily involves a dynamic simulation of the vehicle being controlled, but the accuracy required of that simulation may vary greatly from one application to another. As long as the attitude-control accuracy requirements are low, and the vehicle is relatively rigid, the "dynamics block" in a control system block diagram is easily generated. Modern space vehicles are far from rigid, however, and attitude-control accuracy requirements are increasingly stringent, particularly for optical observations from space vehicles. Further improvements in control system performance depend in many instances on improved simulation of the vehicle dynamics. The importance of this subject is reflected in the existence of the NASA Space Vehicle Design Criteria Monograph, Effects of Structural Flexibility on Spacecraft Control Systems (Ref. 1). This document includes relevant case histories, references, and design recommendations.

The incorporation of vehicle flexibility into control system design is a requirement neither new nor unique to space vehicle applications. Designers of control systems for missiles and aircraft have long ago found it necessary to consider vehicle flexibility in control system development. To some degree, the concepts developed in this earlier context can be applied intact to space applications (Refs. 2-10), but in many respects space vehicles are unique, and new concepts must be developed to deal with them.

The environmental history of every artificial satellite or spacecraft is marked by a brief interval of vigorous acceleration and vibration during boost, followed by prolonged functioning in a quiescent mode of operation characterized by extremely small loads and accelerations. Some missions may require a second period of violent acceleration for all or part of the system, but still there are two distinct and radically different dynamic environments to be considered. The universal solution to the dilemma this poses for the structural designer has been the adoption of lightweight (and extremely flexible) deployable appendages. The resulting vehicle is relatively compact and rigid during the launch phase of its history, but after boost termination it emerges like a butterfly from its cocoon, extending antennas and booms and unfurling solar-cell arrays until the structure has undergone complete metamorphosis.

*The author is an Associate Professor in the School of Engineering and Applied Science at the University of California, Los Angeles.
Although modern space vehicles can be found in a wide variety of configurations, certain dynamic features are sufficiently common to be described as characteristic of space vehicles, as opposed to missiles, aircraft, or land and sea vehicles. Most current space vehicles can be described as the combination of one or more essentially rigid bodies with one or more extremely flexible bodies. This natural separation of the structural subsystems of the vehicles into two distinct regimes is a consequence of the widespread use of deployable appendages, which are much less commonly found on surface or airborne vehicles.

The structural subsystems of a spacecraft or satellite are often required to undergo substantial relative motions during mission performance, while large antennas, solar-cell arrays, instrument packages, or propulsion devices change their relative orientation. With the exceptions of variable-sweep and rotary-wing aircraft, most surface and air vehicles have dynamically significant moving parts only as semirigid rotors in the form of wheels, propellers, and propulsion subsystems. A rigid, symmetric, fixed-axis rotor is easily incorporated into a dynamic simulation of a vehicle, but the presence of a fully articulated flexible antenna on a space vehicle necessitates major changes in the formulation of its equations of motion, particularly when the relative motion of vehicle and antenna is subject to closed-loop, nonlinear control.

A different class of problem is introduced by the use of discrete damping devices in spacecraft subsystem vibration isolation or passive spin-stabilization nutation attenuation.

All of these characteristic features of space vehicles present problems in dynamic simulation, and some of these problems are quite difficult to resolve by application of the methodology of aircraft and missile control system design. There must be a comprehensive examination of the question of dynamic simulation for attitude control of space vehicles, and new approaches must be developed for applications that are beyond the scope of the efficient utilization of traditional methods of analysis.

B. Scope

The present study is concerned with the development of methods for the dynamic simulation of flexible spacecraft. The emphasis in this report is on analytic simulations, although it is recognized that analytic results must generally be confirmed or corrected by experimental measurements on models or prototype vehicles.

Analytic methods treated here are restricted in their applicability to vehicles admitting of idealization as combinations of rigid bodies, particles, continuous elastic bodies, and (in special cases) fluids. The equations of motion of a continuous mechanical system (with continuous spatial variation of mass and flexibility) are partial differential equations. It is assumed in this report that any such equations descriptive of linearly elastic solid subsystems have been subjected to a "modal coordinate" transformation (as defined later in this section) and the resulting coordinates have been truncated to permit representation of system deformation with a finite number of modal coordinates. Thus continuous linearly elastic solids are admissible only in the sense that they can be represented either as a collection of interconnected rigid bodies or by a finite number of modal deformation coordinates. Fluids are admissible under three conditions only: (1) the fluid in a given container can be idealized as rigid, (2) the fluid can be represented by a finite number of modal coordinates, or (3) all aspects of the fluid dynamics can be ignored, except possibly the influence of the fluid on energy dissipation. In consequence of the formal exclusion of continuous mechanical elements, the equations of motion are ordinary (and not partial) differential equations.

Three distinct approaches to the analysis of flexible vehicles are treated in this report: (1) the discrete-coordinate formulation, (2) the hybrid-coordinate formulation, and (3) the vehicle normal-coordinate formulation. These methods are considered in turn in Sections II, III, and IV, with emphasis on the second method. A brief, qualitative description of each of the three general approaches is given in the following paragraphs.

The vehicle normal-coordinate method is the traditional approach to the vibration analysis of elastic systems. This method is well documented in texts (Refs. 11–13), and space applications can be found in technical reports and journal papers (Refs. 2–10), so in this report it is described only insofar as this seems necessary for completeness of the study and for comparison with less familiar methods.

In brief, this method consists of formulating equations of motion, whenever possible, as a system of independent (uncoupled) scalar second-order differential equations. For limited motions of a restricted class of physical systems, systematic procedures can be generated for the determination of the transformation necessary to change from some arbitrarily selected coordinate system to the coordinate system corresponding to uncoupled scalar equations of motion. In general, the new uncoupled co-

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ordinates (called normal coordinates) do not correspond individually to the translations and rotations of discrete points or rigid bodies of the vehicle; instead, each coordinate is associated with a motion in which the entire vehicle participates. Coordinates that correspond to motion of more than one particle or rigid body of the system are called distributed or modal coordinates, as opposed to discrete coordinates. When all the dynamic equations of a system are uncoupled (as in the vehicle normal-coordinate method), the modal coordinates are called normal-mode coordinates for that system. Because the equations of motion are uncoupled, the vehicle can undergo motion in which only one of the scalar coordinates of the system participates. In this mode of motion, all points of the vehicle oscillate at the same frequency (the normal-mode frequency) and the vehicle undergoes periodic deformations into the same deformed shape (the normal-mode shape). Each modal coordinate defines the amount of the response in the corresponding mode, suitably normalized. The independence of these normal-mode coordinates permits the independent calculation of their participation in the vehicle motion. This is the key feature of the normal-coordinate approach, because it permits the exercise of engineering judgment in determining which coordinates are so significant as to warrant retention, and which may be abandoned in coordinate truncation.

Even this classical approach to vibration analysis has not been very widely used in spacecraft control system design and evaluation, so aspects of the method peculiar to this area of application are emphasized in this report. The assumptions underlying the vehicle normal-coordinate approach, and the theoretical and practical limitations on its use, are brought into focus. Appreciation of the restrictions implicit in this approach is particularly important in space vehicle application, because the method breaks down when the system includes nonlinearities, rotors, discrete dampers, or articulated moving parts, and these are precisely the features just described as characteristic of space vehicles. Formulating equations of motion as first-order (state) equations eliminates some of these obstacles, but the practice of working with normal-mode coordinates for the entire vehicle is still quite restricted.

All of these difficulties can be accommodated by avoiding modal coordinates entirely, relying upon a complete set of equations of a collection of interconnected rigid bodies considered to be representative of the vehicle. Because each rigid body of the system model is discrete, and the coordinates employed in this approach are coordinates of position and/or attitude of the individual bodies, this approach is sometimes called the discrete-parameter or discrete-coordinate method.

The discrete-parameter approach to space vehicle simulation has received great emphasis since 1965, primarily because of its generality. A growing body of literature on this subject is becoming available (Refs. 14–22), and digital computer programs based on these and other formulations are becoming commonplace tools of analysis.

The approach most frequently adopted (Refs. 14–21) involves direct application of the Newton-Euler equations of translation and rotation to various subsets of bodies in the assembly (often to each of the bodies individually). Alternatively, Lagrange's equations are sometimes applied (Ref. 22).

When application is restricted to a vehicle model composed of \( n \) rigid bodies interconnected at \( n - 1 \) point contacts so as to form a “tree” structure topologically, the Newton-Euler equations allow dramatic simplification. These restrictive assumptions are illustrated in Fig. 1, which shows that adjacent rigid bodies share at least one common point, and no closed loops are formed by any string of rigid bodies. The 12 bodies and 11 points

Fig. 1. Topological tree of point-connected rigid bodies
of contact in the figure are arbitrarily labeled. The simulations described in Refs. 14–19 and 21 are limited in this manner (the "tree" of Ref. 21 is more like a "bush," with a trunk and limbs each consisting of a single rigid body). Under these conditions, for a complete kinematic description of the system, it is sufficient to know the location of one of its points (e.g., the mass center) and the orientation or attitude of each of the bodies. It is therefore possible to reduce the number of equations required for the dynamic simulation, eliminating all translational coordinates (except those of the system mass center, which may be determined independently in many cases).

Euler's equations of rotational motion of a rigid body (say, the $i$th body) are three scalar first-order differential equations in the variables $w_i, w_j, w_k$, which are measure numbers of the inertial angular velocity of that body for a body-fixed vector basis. These variables, collected for all $n$ bodies, generally constitute the unknowns of the final differential equations, which are therefore first-order equations. Because a set of angular velocity measure numbers does not in itself provide a complete kinematic description of the system, an additional set of first-order equations must be included to permit the determination of attitude by angular velocity integration. These kinematic equations may be expressed in terms of any desired set of attitude parameters, e.g., direction cosines or a subset of direction cosines, Euler parameters, Euler-Rodrigues parameters (Ref. 23), Euler angles (3, 1, 3 rotations), or Tait-Bryan angles (1, 2, 3 rotations). The last three alternatives involve only three parameters (the minimum number) for the attitude description of each body, but all three-parameter sets are plagued by isolated singularities that make numerical computations impossible for certain attitudes (without coordinate transformation). Most spacecraft-simulation programs employ a larger set of attitude parameters, accepting the inconvenience of working with a set of coordinates interrelated by one or more constraint relationships. There appears to be no overwhelming preference between a subset of the direction cosines (usually six in number) and the four Euler parameters (or Cayley-Klein parameters or elements of a quaternion).

If one of the bodies of the set has more than one point in common with another body (e.g., the two are connected by a line hinge), this may be represented in the simulation by an additional constraint equation. Procedures exist for the systematic incorporation of such "joint constraints" into the dynamic equations, with consequent reduction in dimension of the matrix differential equation of the system. Analysts disagree, however, on the advisability of this operation for computational efficiency (Refs. 14, 15, and 20).

When Lagrange's equations are written in terms of a set of independent generalized coordinates (Ref. 24, article 26), constraint equations are absorbed automatically, and the equations of motion are obtained immediately as a set of second-order differential equations of minimum number. The restriction to the use of independent generalized coordinates may, however, be a handicap, since this precludes the adoption of Euler parameters or direction cosines for attitude description. As a result, such a conventional Lagrangian formulation must employ a coordinate system not entirely free of singularities. In practice, most (perhaps all) programs based on Lagrange's equations use attitude angles as coordinates (rather than Euler-Rodrigues parameters), and this selection introduces computationally cumbersome trigonometric functions into the calculations. This may be a greater handicap than the presence of singularities, particularly if digital simulation is anticipated. This approach is also restricted to systems with holonomic constraints, although this is perhaps not as severe a restriction in aerospace applications as it would be for land vehicle applications that include rolling wheels.

One may increase the scope and flexibility of the Lagrangian approach in at least two significant ways. With the introduction of Lagrange multipliers (Ref. 24, article 87), the restriction to a set of independent generalized coordinates is relaxed, and one may adopt any kinematically complete set of coordinates, providing only that constraint equations in the form of equalities (not inequalities) exist in sufficient number to offset the coordinate redundancy. (Unfortunately, aerospace vehicle subsystem constraints such as damper and gimbal "stops" are in the excluded class of inequalities.) This approach extends the scope to certain nonholonomic systems, and it permits the use of redundant attitude parameters (e.g., Euler parameters). The price paid is the added dimension of the problem. This method does not seem promising in comparison with the Newton-Euler method, and appears not to have been pursued for general space vehicle simulation. However, applications to restricted problems can be found in the aerospace journal literature (Ref. 25).

The second relevant modification of the Lagrangian formulation is more fundamental. It is possible to avoid the reliance in Lagrange's equations upon generalized coordinates, which must be sufficient to establish fully in
themselves the configuration of the system at any given time. One may alternatively write Lagrange's equations in terms of quasi-coordinates (Ref. 24, article 30), which are quantities whose differentials may be written as linear combinations of differentials of generalized coordinates and time. Since the angular velocity measure numbers \( w_i^1, w_i^2, w_i^3 \) of a given \( i \)th body qualify as derivatives of quasi-coordinates, this Lagrangian approach can produce results with some of the qualities of the Newton-Euler formulation.

A recent adaptation by Kane and Wang (Refs. 26, 27) of the quasi-coordinate formulation seems well suited to complex spacecraft simulation. The method is applicable to certain nonholonomic systems and to systems with redundant (and constrained) coordinates; yet it automatically eliminates nonworking constraint forces and torques. Although the approach by Kane and Wang appears to combine certain advantages of the Newton-Euler method and the Lagrangian method, this path to complex system simulation has not yet been taken to the point of a multipurpose computer program for machine computation of the response of arbitrary discrete parameter systems.

Section II contains an introduction to the formulation of the equations of motion for discrete-parameter systems. The text describes without derivation the results of Refs. 14 and 15, and notes some of the features of Refs. 16–20. The method of Kane and Wang is also briefly described. In view of the functional and structural beauty of the equations developed in these references, it seems unlikely that improvement can be realized by additional independent derivations, and the analyst confronting the problem of simulating complex discrete-parameter systems is advised to acquaint himself first with the referenced literature.

Discrete-parameter simulations are not without serious disadvantages. Satisfactory simulation of real vehicles may require a great many rigid bodies in the model. The resulting differential equations are then of high dimension, and their digital solution may be plagued by high-frequency responses that are of no interest to the engineer. There is no mechanism for truncating the matrix of coordinates retained in the determination of vehicle response, so as to eliminate these high-frequency responses and reduce the dimension of the problem.

When the system configuration demands the generality of a discrete-parameter simulation, this alternative must be adopted. But when all or part of the vehicle allows the assumption of small, linearly elastic deformation, the most efficient simulation is that which combines discrete coordinates with distributed (modal) coordinates, retaining the generality of discrete coordinates where necessary, but securing the computational advantages of (truncated) modal coordinates where possible. The result is a hybrid-coordinate system that permits accurate simulation of complex modern space vehicles with a minimum number of coordinates, and with irrelevant high-frequency oscillations removed from the integration output by modal-coordinate truncation. Although this method is a natural combination of the other two approaches, it has received very little emphasis in the technical literature to date (Refs. 28, 29). For this reason, and because this method seems to combine the computational advantages of modal analysis with the needed generality of the discrete-coordinate approach, the hybrid-coordinate method receives major emphasis in this report.

The question of control system simulation per se is included in this report only insofar as this issue is coupled to the primary question of flexible vehicle simulation.

II. Discrete-Coordinate Methods

A. Augmented-Body Methods

Hooker and Margulies (Ref. 14) and Roberson and Wittenberg (Refs. 15, 16) have observed that when a system of \( n \) point-connected rigid bodies is assembled in a topological tree (as in Fig. 1), certain inertia-like terms naturally appear in combination in the individual equations of motion of each of the rigid bodies in the set. These combinations admit of physical interpretation as the inertia dyadics (or tensors or matrices) of abstractions called the augmented bodies. Briefly, the \( i \)th augmented body consists of the \( i \)th body of the set together with certain particles (point masses) attached to each of the joints of that body. The point mass attached to a given joint of the \( i \)th body equals the total mass of all of the connected bodies located "outboard" of the joint. For example, the augmented body 9 of the 12-body system shown in Fig. 1 is illustrated in Fig. 2. The mass center of the augmented body is called the connection barycenter (or simply the barycenter). The inertia dyadic of the augmented body with respect to the corresponding barycenter is the term that appears in the equations.

A variety of approaches might be considered in formulating the Newton-Euler equations of motion of a system of rigid bodies such as illustrated in Fig. 1. Perhaps the most direct approach is to isolate each of the bodies in
turn and write the translational and rotational vector equations for that body, recognizing that neighboring bodies contribute to the applied force and torque. This procedure leads to the appearance of augmented-body inertia dyadics, so it is the method used here. It should be recognized, however, that this is not the only possible approach; one might, for example, exclude the equations of motion of the nth body and instead write the translational and rotational equations of motion of the total system, or of any connected group of bodies within the system. Other possibilities are explored in Section II-B.

For an individual body, say, the ith body, the translational and rotational vector equations of motion may be written as

\[ \begin{align*}
F^i &= m_i a^i \\
T^i &= H^i = \dot{H}^i = I^i \cdot \dot{a}^i + \omega^i \times I^i \cdot \omega^i
\end{align*} \tag{1} \]

where \( F^i \) is the resultant force and \( T^i \) the resultant torque applied to the ith body, \( m_i \) is the body mass, \( a^i \) the mass center inertial acceleration, \( H^i \) the body angular momentum, \( I^i \) the inertia dyadic, and \( \omega^i \) the inertial angular velocity. The quantities \( T^i, H^i, \) and \( I^i \) are all referred to the body mass center. In the notation used in this report, dot (\( \dot{\cdot} \)) over a vector indicates time differentiation in an inertial reference frame.

Among the constituents of the applied force \( F^i \) there are “hinge forces,” i.e., forces applied to the ith body by contiguous bodies at the points of contact. These hinge forces generally contribute also to the torque \( T^i \) about the ith body mass center, and typically this is the only coupling mechanism between the translational and rotational equations (1).

It is evident by inspection of Fig. 1 that knowledge of the attitude of every body of a point-connected set is sufficient for the complete determination of the system configuration and attitude; it is not necessary to know mass-center position coordinates as well, since these must follow from the kinematic constraints imposed by the joints. Consequently it must be possible to combine the vector equations (1) for all \( n \) bodies in such a way as to obtain \( n \) vector equations of rotational motion that are internally complete, without any coupling terms to additional translational equations. This is accomplished by first solving the translational equations (1) for the hinge forces in terms of mass-center accelerations, and then obtaining from kinematics the mass-center accelerations in terms of geometry and rotational coordinates.

The specific manipulations that accomplish this reduction of equations (1) to a dynamically complete set of half as many equations can be found in Refs. 14-18. In the work of Hooker and Margulies (Ref. 14), the manipulation is performed at the level of explicit vector equations for the individual bodies, while in the work of Roberson and Wittenberg (Refs. 15, 16), this task is deferred to a stage of the derivation in which the rotational equations of motion are matrix equations of dimension \( 3n \). In both developments, there emerge terms (dyadics in Ref. 14 and matrices in Refs. 15, 16) that have the indicated interpretation as inertia dyadics or matrices of augmented bodies. The vector-dyadic result, for example, is given for the \( \lambda \)th body in the notation of Ref. 14 by

\[ \Phi^\lambda = \omega_\lambda + \omega_\mu \times \Phi^\mu \]

\[ T^\lambda = T^\lambda'' + \sum_{j \in J_\lambda} T_{\lambda j}'' + \sum_{\mu \neq \lambda} D_{\mu \lambda} \times F_\mu + \sum_{\mu \neq \lambda} D_{\mu \lambda} \times \left[ \omega_\mu \times D_{\mu \lambda} + \omega_\mu \times \left( \omega_\mu \times D_{\mu \lambda} \right) \right] \]

with the following symbol definitions:

\( \Phi^\lambda \) = the inertia dyadic of the \( \lambda \)th augmented body, referred to the corresponding barycenter

\( \omega_\lambda \) and \( \omega_\mu \) = the inertial angular velocities of bodies \( \lambda \) and \( \mu \), respectively

\( T^\lambda \) = that portion of the resultant torque applied to body \( \lambda \) obtained by excluding forces and torques applied at joints

\( T_{\lambda j}'' \) = the “hinge torque” applied at joint \( j \) of body \( \lambda \)

\( J_\lambda \) = the set of numeric labels for the joints on body \( \lambda \) (e.g., from Figs. 1 and 2, \( J_\lambda \) includes 7, 8, and 9)
\[ D_\lambda = \text{the position vector from the barycenter } B_\lambda \text{ to the mass center of body } \lambda \text{ (e.g., see Fig. 2)} \]

\[ F_\lambda \text{ and } F_\mu = \text{the forces applied to bodies } \lambda \text{ and } \mu, \text{ respectively, excluding forces applied at joints} \]

\[ D_{\lambda\mu} = \text{the position vector from the barycenter } B_\lambda \text{ to the joint of body } \lambda \text{ that leads to body } \mu \text{ (even if body } \mu \text{ is not directly connected to body } \lambda, \text{ but instead is part of a chain of bodies connected to body } \lambda) \]

\[ \mathcal{Q}_n = \text{the total system mass} \]

\[ \sum_{j \in \Lambda} = \text{the sum over values of } j \text{ in the set } \Lambda. \]

The left side of Eq. (2) has exactly the form of the vector-dyadic representation of Euler’s equations, except for the substitution of the augmented-body inertia-dyadic about the barycenter \( \Phi^i \) for the body inertia-dyadic about the mass center \( \mathbf{I}^i \). The right side of Eq. (2) includes the body torque \( T_{\lambda} \) and the relevant hinge torques \( T_{\mu j} \), as would be the case for Euler’s equations, but the torque contribution of hinge forces takes a surprising form. In every case, torques are calculated with respect to the barycenter, and the appropriate force turns out to be the sum of the body forces \( F_\mu \) and certain “inertial forces” that may be attributed to angular accelerations and centripetal accelerations.

In view of the complexity of the n-body system under examination, it is astonishing that the equations of motion should be so simple in structure and amenable to physical interpretation. Simplification is even more dramatic when there is a coincidence of barycenter \( B_\lambda \) and a joint, since then the vector \( D_{\lambda\mu} \) is zero for all bodies connected to body \( \lambda \) by that joint, and substantial decoupling of the equations results.

In general, of course, there are required \( n \) vector-dyadic equations such as Eq. (2) to complete the dynamic description, and in addition some specification of the hinge torques \( T_{\mu j} \) is required. In most applications, some of the bodies in the model are connected by line hinges, so that a line is common to two bodies of the system. The hinge torques \( T_{\mu j} \) transverse to the hinge axis then become constraint torques, and each line hinge provides two scalar constraint equations that preclude relative rotations except about the hinge axis. Similarly, a two-gimbal joint provides one constraint equation. If there are \( v \) constraint equations and \( n \) rigid bodies in the system, then there are available \( 3n + v \) scalar equations to be solved for the \( 3n \) unknown angular velocity measure numbers plus the \( v \) constraint torques. (An additional equation is of course required for the determination of the translation of the system mass center, but this equation is generally uncoupled with the attitude equations and is ignored here.)

The most direct approach to the determination of dynamic response is to combine all the equations of dynamics and kinematics with the constraint equations as a single-matrix first-order differential equation of dimension \( 6n + v \) and proceed with numerical integration.

The possibility of eliminating the \( v \) constraint torques, thus reducing the dimension of the problem to \( 6n \), is discussed briefly in Ref. 14, and related questions are explored extensively in Refs. 16, 18, 19, 20, 26, and 27. Some of the methods that serve this purpose are discussed in Sections II-B and II-C.

The task of assembling the \( n \) vector-dyadic rotational equations into a single matrix equation is made quite awkward by the summations over limited sets of joints and bodies that appear in Eq. (2). Roberson and Wittenberg (Refs. 15, 16), in a derivation that developed in parallel to that of Hooker and Margulies (Ref. 14), utilize graph theory to construct an elegant formalism for the systematic assemblage of the individual vector-dyadic equations into the single \( 3n \)-dimensional matrix equation of motion of the total system. The equations of Ref. 15 are programmed (with some modification) in Ref. 17, and those of Ref. 14 provide the basis for the program developed in Ref. 18.

Whether the matrix equation is developed with the beautiful formalism devised by Roberson and Wittenberg or assembled in more pedestrian fashion (as computational efficiency may suggest), the final structure of the equation must be as follows:

\[ \mathcal{R} \dot{\omega} = \mathcal{B} + \mathcal{L} \quad (3) \]

where \( \mathcal{L} \) absorbs all terms involving constraint torques, \( \mathcal{B} \) embraces all external forces and torques as well as all terms involving inertial angular velocity terms \( \omega_s^j (\alpha = 1, 2, 3, i = 1, \cdots, n) \), and \( \mathcal{R} \) is the coefficient matrix of all differentiated terms, which are assembled in the column matrix \( \dot{\omega} \). Ignoring for the moment the question of how constraint equations are to be used to accommodate unknowns in \( \mathcal{L} \), one can readily see that
it is the nature of $\mathcal{A}$ that will determine computational
difficulty. If $\mathcal{A}$ were a nonsingular diagonal matrix, a
trivial inversion would permit Eq. (3) to be written as
\[ \dot{\mathcal{A}} = \mathcal{A}^{-1} \mathcal{B} + \mathcal{A}^{-1} \mathcal{L} \] (4)
where superscript $-1$ denotes matrix inverse.

Unfortunately, a glance at Eq. (2) indicates that iner-
tial coupling terms involving $\dot{\mathcal{A}}_\mu$ must exist unless $\mathcal{D}_{\lambda\mu} = 0$
for all $\mu$. The same term $\mathcal{D}_{\lambda\mu} \times \dot{\mathcal{A}}_\mu$ also assures the time-
varying character of $\mathcal{A}$, since the cross product depends
on the changing relative attitude of bodies $\lambda$ and $\mu$. This
constitutes a major obstacle to numerical integration, since
if Eq. (4) is used it becomes necessary to invert the $3n$ by $3n$
matrix $\mathcal{A}$ at each step of the integration. In practice, it may prove more convenient to retain the
equation in the form of Eq. (3), applying Gaussian elimina-
tion rather than matrix inversion at each integration step.
Even with this expedient, the algebraic process of finding $\dot{\mathcal{A}}$ is apt to consume most of the computer time
in numerical integration. Since time required for this process increases roughly with the third power of the
dimension of the matrix $\mathcal{A}$, abundant motivation exists
for working with equations of minimum dimension.

B. Nested-Body Methods

In writing $n$ vector equations of rotational motion for
a set of $n$ point-connected rigid bodies, one must make a
choice of the $n$ material subsystems to be isolated for
dynamic analysis; it is not obviously advantageous to isolate each of the rigid bodies individually, as is gen-
erally the practice in Refs. 14-18. (In Refs. 15-18 there
is an explicit departure from this practice when a pair
of rigid bodies consists of a rigid body containing a rigid,
axisymmetric rotor on a fixed axis.)

Velman (Ref. 20) and Russell (Ref. 19) elect instead to
write vector equations of motion in turn for $n$ different
subsets of bodies, including a final set of rotational and
translational equations for the composite vehicle. For
example, if the vehicle were modeled by 12 point-
connected rigid bodies as shown in Fig. 1, both Velman
and Russell might write equations of motion first for
body 12, then for the subset including bodies 11 and 12,
and then for 10, 11, 12. Next, body 8 might be isolated
and its equations written, and these followed by equa-
tions for the subset including bodies 8-12. One might
then begin anew at the end of another chain of bodies
(either body 1 or body 7), and progress inward as far
as possible without ever including more than one addi-
tional body in any one step, and without ever considering
a subset with more than one joint connecting it with the
excluded bodies of the system. For this example, one
might follow the indicated pattern until each of the three
chains attached to body 4 has been considered, without
ever including body 4 itself. (This choice of body 4 is
arbitrary, since other subset selections within the pre-
scribed pattern can converge as well on any other body
of the system.) Finally, the equations of motion of the
composite vehicle are recorded, to complete the dynamic
simulation.

The idea of isolating in sequence such subsets of rigid
bodies in an $n$-body system (called nested bodies by Velman) seems to have both advantages and disadvan-
tages. The concepts of the augmented body and the
connection barycenter are helpful aids to physical inter-
pretation (if not to numerical computation), and the
terms with which these phrases are identified in the
equations of Roberson-Wittenberg (Refs. 15, 16) and
Hooker-Margulies (Ref. 14) do not appear when the
nested-body approach is taken. In compensation, how-
ever, it would appear that the nested-body derivation
facilitates the elimination of internal constraint forces
and torques.

Velman (Ref. 20) derives his equations in a manner
that leads quite naturally to the use of attitude variables
that establish the orientation of each body of the system
relative to an adjacent body (except of course for one
reference body with inertial attitude parameters). Because
geometrical constraints generally restrict relative motions,
it is perhaps easier to identify which attitude variables
are constrained when relative motion coordinates are
employed. Nonetheless, there is no attempt to eliminate
kinematically constrained variables in the derivation of
Velman’s dynamical equations, so his equations have at
the outset the same basic structure and dimension as those
of Roberson and Wittenberg or Hooker and Margulies; namely,

\[ \mathcal{A} \dot{\mathcal{A}} = \mathcal{B} + \mathcal{L} \] (5)

These symbols differ in interpretation among the several
authors, and, in particular, Velman’s $\dot{\mathcal{A}}$ includes some rela-
tive velocities. It should be noted also that Velman’s sim-
ulation specifically includes a linearly oscillating particle,
so the method is not restricted to a point-connected set
of rigid bodies. In Eq. (5), then, one of the scalars in the
matrix $\mathcal{A}$ is the translational velocity of a particle relative
to its housing body. As in Refs. 14-16, the dimension of
matrix $\mathcal{A}$ is $3n$ for an $n$-body system as long as each body
is either a particle or a rigid body forming part of a point-connected set. Again, \( \mathcal{A} \) and \( \mathcal{B} \) are matrices that depend upon the kinematic unknowns of the problem, and \( \mathcal{L} \) is the matrix of unknown constraint torques (and constraint forces for the oscillating particle). The most significant feature of Velman's work is the procedure he devises for the elimination from these equations of the unknown constraint forces and torques.

Velman notes that the effect of the constraints is usually to confine the solution of Eq. (5) to some linear manifold in the \( 3n \)-dimensional space of \( \omega \). If, for example, a line hinge connects two rigid bodies of the system, and three of the scalars in \( \omega \) are the relative rotation rates of these bodies about the hinge axis and two transverse axes, the effect of the constraints is to confine the solution to that subspace of the \( \omega \) space excluding the two transverse-axis rotation rates, which are constrained by the hinge to be zero. The subspace to which the solution is confined is then of dimension \( 3n - 2 \). This fact is unchanged even when the choice of variables in \( \omega \) does not specifically include these three relative rates about and transverse to the hinge; this choice may simplify the explicit specification of the solution subspace, but it introduces no conceptual change in the argument. Similarly, the line of argument remains intact when the solution in a subspace of \( \omega \) is constrained to be not zero, but a specified function of time. If, for the previous example, the two bodies connected by a line hinge undergo a prescribed relative rotation about that hinge (as for an idealized scanning antenna), then the solution is a prescribed function of time in a three-dimensional subspace of the \( \omega \) space.

The preceding remarks amount to the observation that the constraints determine the projection of the motion on a particular linear manifold in the \( \omega \) space. Velman introduces the perpendicular projection operator \( \mathcal{D} \) (a \( 3n \times 3n \) matrix called \( E \) in Ref. 20) and the analytic expression

\[
\mathcal{D} \dot{\omega} = \mu(t)
\]

and the kinematic constraint equation would be

\[
\mathcal{D} \dot{\omega} = [0 \, 0 \, \dot{\omega}_4 \, \dot{\omega}_5 \, \dot{\omega}_6]^{T}
\]

\[
= [0 \, 0 \, 0 \, 0 \, \ddot{\theta}(t)]^{T}
\]

where superscript \( T \) denotes the matrix transpose.

Rather than augment the \( 3n \) scalar dynamic equations in Eq. (5) by the \( r \) scalar constraint equations that constitute the nontrivial part of Eq. (6), Velman (Ref. 20) uses the latter to eliminate the constraint torques \( \mathcal{L} \) from the former. To accomplish this, he notes that the vector of constraint torques (and forces) \( \mathcal{L} \) lies in the same manifold of the \( \omega \) space in which motion is prescribed. Thus if \( \mathcal{D} \) is defined as the supplementary perpendicular projection operator for \( \mathcal{D} \), so that \( \mathcal{D} + \mathcal{D} \) is the identity matrix \( E \),

\[
\mathcal{D} + \mathcal{D} = E
\]

then \( \mathcal{D} \) operating on \( \mathcal{L} \) is given by

\[
\mathcal{D} \mathcal{L} = 0
\]

For the simple example of the two hinged bodies, Eq. (10) appears in expansion as

\[
[1 \, 0 \, 0 \, 0 \, 0 \, 0] [0] = [0]
\]

\[
[0 \, 1 \, 0 \, 0 \, 0 \, 0] [0] = [0]
\]

\[
[0 \, 0 \, 1 \, 0 \, 0 \, 0] [0] = [0]
\]

\[
[0 \, 0 \, 0 \, 0 \, 0 \, 0] [L_4] = [0]
\]

\[
[0 \, 0 \, 0 \, 0 \, 0 \, 0] [L_5] = [0]
\]

Now one can return to Eq. (5) and rewrite it as

\[
\mathcal{A} (\mathcal{D} + \mathcal{D}) \dot{\omega} = \mathcal{B} + \mathcal{L}
\]
which upon premultiplication by $\mathcal{J}$ and substitution of Eqs. (6) and (10) becomes

$$\mathcal{J}\mathcal{A}\dot{\mathcal{J}} = \mathcal{J}\mathcal{B} - \mathcal{J}\mathcal{A}\mu(t)$$

(13)

Fleischer notes that the matrix $\mathcal{J}\mathcal{A}\mathcal{J}$ is singular, so Eq. (13) is not yet ready for machine computation. Using the idempotency of the perpendicular projection operators and their supplementary character, one may write

$$\mathcal{J}\mathcal{P} = \mathcal{P}, \quad \mathcal{J}\mathcal{J} = \mathcal{J}, \quad \mathcal{J}\mathcal{P} = 0$$

(14)

and then rewrite Eq. (13) in the form

$$\mathcal{J}\mathcal{A}\dot{\mathcal{J}} = \mathcal{J}\mathcal{B} - \mathcal{J}\mathcal{A}\mu(t)$$

(15)

The matrix $(\mathcal{J}\mathcal{A}\mathcal{J} + \mathcal{P})$ can be inverted to obtain

$$\mathcal{J}\dot{\omega} = (\mathcal{J}\mathcal{A}\mathcal{J} + \mathcal{P})^{-1} \mathcal{J}[\mathcal{B} - \mathcal{A}\mu(t)]$$

(16)

which when added to Eq. (6) yields

$$\dot{\omega} = (\mathcal{J}\mathcal{A}\mathcal{J} + \mathcal{P})^{-1} \mathcal{J}[\mathcal{B} - \mathcal{A}\mu(t)] + \mu(t)$$

(17)

since $\mathcal{J} + \mathcal{P} = \mathcal{E}$ from Eq. (9). This is the form of the equation used by Velman. The dimension of the unknown matrix $\dot{\omega}$ is still $3n$, but the constraint torques have been eliminated.

As noted earlier, the fact that Velman's variables in $\omega$ are generally relative motions simplifies the physical interpretation of the constraint equation (6), and often results in constant perpendicular projection operators $\mathcal{P}$ and $\mathcal{J}$. Still, the coefficient matrix $\mathcal{A}$ depends on the equation variables and accordingly varies with time, so the $3n$ by $3n$ matrix $(\mathcal{J}\mathcal{A}\mathcal{J} + \mathcal{P})$ must be inverted at each integration step for the dynamic equation (17). Since the system has only $3n - v$ degrees of freedom, one might hope to reduce the dimension of the variable $\dot{\omega}$ in Eq. (17), and in the process reduce the size of the matrix requiring inversion at each integration step.

Fleischer (Ref. 18) has developed a procedure for accomplishing the indicated reduction of the dimension of the problem to $3n - v$. If the variable $\omega$ is partitioned as $\omega' = [\omega_f \omega_c]^T$, with subscripts $f$ and $c$ identifying free and constrained coordinates, then one may attempt to use the constraint equations to find $\omega_c$ in terms of $\omega_f$, and then retain only that portion of Eq. (17) involving $\dot{\omega}_f$ on the left side (i.e., truncate the matrix $\dot{\omega}$ to $\dot{\omega}_f$). Actually, Fleischer pursues a somewhat different path from the basic equations (5) and (6) to the final equations of motion, and in the process he introduces new operators to facilitate the reduction of dimension of the final equations. Because the appropriate modifications are fully treated in Ref. 18, they are not included here.

Fleischer's final equations have the form

$$\mathcal{A}_f \dot{\omega}_f = \mathcal{B}_f$$

(18)

in which $\mathcal{A}_f$ and $\mathcal{B}_f$ are time-varying and of dimension $(3n - v)$ by $(3n - v)$. Computation requires the inversion of $\mathcal{A}_f$ at each integration step (or the equivalent Gaussian elimination process), and since this inversion is the most time-consuming part of a major simulation program, this reduction of dimension would appear to be a substantial improvement. Yet there are additional computer operations involved in obtaining and solving the reduced equations, and differences in the detailed structure of the equations may impede computation in the reduced case, so perhaps it would be useful to discuss the relative computational advantages of the several forms of the discrete-parameter system equations. For example, buried within the matrices $\mathcal{A}_f$ and $\mathcal{B}_f$ in Eq. (18) there is the inverse of a matrix of dimension $v$. When $\omega$ comprises relative motion variables (as in Velman's derivation), the matrix to be inverted is generally constant, requiring only one inversion operation. But when $\omega$ comprises inertial rates of the various bodies, this matrix depends on time, and the matrix inverse must be obtained with each integration step. In Fleischer's work (Ref. 18), the basic dynamic equations of Hooker and Margulies (Ref. 14) are combined with a modified version of Velman's constraint elimination procedure (Ref. 20), so inertial rates are the variables and the indicated repeated inversion is required. At each step of the integration, Fleischer must invert $\mathcal{B}_f$ by $v$ matrix and then invert (or apply Gaussian elimination to) a matrix of dimension $(3n - v)$ by $(3n - v)$, whereas Velman must invert (or apply Gaussian elimination to) one matrix of dimension $3n$ by $3n$. As noted, Hooker and Margulies appear to favor working directly with the dynamic equations as augmented by the constraint equations, accepting system equations of dimension $(3n + v)$ by $(3n + v)$. It seems probable that selection of an optimum approach will vary from one application to the next, but that for very complex systems the advantage would lie with the approach yielding equations of the lowest dimension.

In the derivations of Refs. 14–18 and 20, the approach has been first to write dynamic equations in which constraint forces and torques appear, and then (in some
cases) to use constraint equations to eliminate from the dynamic equations the constraint forces and torques and the redundant kinematic variables. This elimination is not accomplished by the analyst before computation, but in general is accomplished by the computer during numerical computation.

Other approaches have been devised that avoid constraint forces and torques from the outset, and thereby preclude the necessity of using computer time to eliminate constraint forces and torques and reduce the number of scalar equations to the number of degrees of freedom. In the context of the Newton-Euler formulation, the objective of avoiding constraint forces for a point-connected n-body system in a topological tree has been pursued successfully by Russell (Ref. 19). Kane and Wang (Refs. 26, 27) accomplish this objective even more generally, using a method described in Section II-C. Although neither Russell nor Kane and Wang support the concept of developing a single multipurpose computer program suitable for a wide range of vehicles, Russell does provide in Ref. 19 an explicit procedure for constructing equations for simulation of an n-body system restricted as above.

Russell adopts the nested-body concept advanced by Velman and described in Section II-A, where the 12-body system of Fig. 1 was discussed as an example. Recall that for this example one may first write the vector rotational equations of motion for body 12, then for subsystem 11 plus 12, etc. As pursued by Velman, this path does involve constraint torques.

Following Russell's approach, the analyst must observe that body 12 is connected to body 11 by a line hinge, and then write for body 12 only that component of the rotational equation $\mathbf{T}^{12} = \mathbf{H}^{12}$ paralleling the hinge axis. This avoids the constraint torques (assuming unspecified rotation about this axis), and it yields one scalar equation corresponding to the single degree of freedom of body 12 relative to body 11. Because the torque $\mathbf{T}^{12}$ is measured relative to the mass center of body 12, the interaction force applied to body 12 by body 11 contributes to $\mathbf{T}^{12}$, and this unknown constraint force must be eliminated by use of the translational equation $\mathbf{F}^{12} = m_{12} \mathbf{a}^{12}$ (Ref. 19). The method advanced by Russell retains the components of the inertial angular momenta of the individual bodies as the unknowns, obtaining angular velocity of the ith body as necessary from the matrix relationship

$$\omega^i = (I^i)^{-1} \mathbf{H}^i$$

(19) (For this reason Russell calls his method the momentum approach.) This distinction is central to the computational question, but not critical to the avoidance of constraint torques.

Having obtained a single scalar equation for the component of $\mathbf{H}^{12}$ along the hinge axis, one next proceeds to consider the subsystem of bodies 11 and 12, writing translational and rotational equations as previously. Now the analyst must inspect the single joint between this subsystem and the rest of the vehicle, namely, the joint connecting bodies 11 and 10. If this is a 3-degree-of-freedom joint, the entire vector rotational equation of the subsystem 11 plus 12 is retained, but if this is a rotationally constrained joint, only those components of the vector equation that do not introduce constraint torques are retained. Whatever the nature of the joint, the number of scalar equations added corresponds to the number of degrees of freedom added in considering a new subsystem. Proceeding in this way, following the pattern of selecting subsystems previously described, one can systematically accumulate as many dynamic equations as there are independent unknowns without introducing constraint torques at all. A simple accounting procedure permits the distribution of the angular momentum of any subsystem among its constituent bodies, and from these individual inertial angular momenta the inertial angular velocities follow from Eq. (19). The dynamic equations then combine with kinematic equations just as they do for the alternative Newton-Euler formulations (Refs. 14-18, 20).

C. Generalized-Force Methods

Among the many methods employing generalized forces, most are within the framework of Lagrange's equation

$$\frac{d}{dt} \left( \frac{\partial L}{\partial \dot{q}_i} \right) - \frac{\partial L}{\partial q_i} = Q_i, \quad i = 1, \ldots, n_f$$

(20)

where $q_1, \ldots, q_{n_f}$ are a complete and independent set of generalized coordinates, the Lagrangian $L$ is the difference in kinetic and potential energy, and the generalized force $Q_i$ is defined in terms of applied forces $\mathbf{F}_1, \ldots, \mathbf{F}_n$ and their inertial position vectors $\mathbf{r}_1, \ldots, \mathbf{r}_n$ by

$$Q_i = \sum_{j=1}^{n} \mathbf{F}_j \cdot \frac{\partial \mathbf{r}_j}{\partial q_i}, \quad i = 1, \ldots, n_f$$

(21)

Lagrange's equation in the form of Eq. (20) is too familiar to warrant review and too limited to warrant adoption as the multipurpose procedure for discrete-parameter system simulation. As noted in the Introduction, restriction
to a complete and independent set of generalized coordinates \( q_1, \ldots, q_n \). This is inhibiting in that it precludes the use of redundant, singularity-free attitude variables and excludes nonholonomic systems. (A holonomic system can be fully described by a set of coordinates \( q_1, \ldots, q_N \), which are related by \( n \) equalities of the form \( f_i(q_1, \ldots, q_N, t) = 0 \), \( i = 1, \ldots, n \), thus permitting, at least theoretically, the determination of \( \dot{v} \) coordinates in terms of the remaining set of \( n_i = N - \nu \) coordinates.)

The generalized forces \( Q_i \) are attractive, however, in that the definition in Eq. (21) serves to eliminate from \( Q_i \) any nonworking constraint forces, and thus to eliminate these unknown and unwanted quantities from the equations of motion.

Kane and Wang (Refs. 26, 27) have devised an approach that retains generalized forces with this desirable property, but permits application to certain nonholonomic systems and to systems with a wider range of acceptable coordinates.

Consider a dynamic system described by a complete but redundant set of coordinates \( q_1, \ldots, q_N \) subject to the \( \nu \) constraint equations typified by

\[
\sum_{i=1}^{N} A_{ij} \dot{q}_i + B_j = 0, \quad j = 1, \ldots, \nu
\]  

(22)

where \( A_{ij} \) and \( B_j \) are functions of \( q_1, \ldots, q_N \) and \( t \). If Eq. (22) cannot be integrated to obtain constraint equations in the form of equalities not involving generalized velocities, this is typical of a class of nonholonomic systems.

In application to a system of \( n \) particles and rigid bodies, one may without restriction write the Newton-Euler equations for the \( i \)th body of the system in the form

\[
\begin{align*}
\mathbf{F}^i + \mathbf{F}^i \, ^{\ast} &= 0, \\
\mathbf{T}^i + \mathbf{T}^i \, ^{\ast} &= 0,
\end{align*}
\]  

(23)

\( i = 1, \ldots, n \)

(24)

where

\[
\mathbf{F}^i \, ^{\ast} = -m_i \dot{a}^i
\]

(25)

and

\[
\mathbf{T}^i \, ^{\ast} = -\mathbf{H}^i
\]

(26)

with \( \mathbf{T}^i \), \( a^i \), and \( \mathbf{H}^i \) referenced to the mass center for the rigid bodies. In application to a rigid body, \( \mathbf{F}^i \) is the resultants applied force. Because the constraints imposed in Eq. (21) imply the presence of constraint forces in \( \mathbf{F}^i \) and constraint torques in \( \mathbf{T}^i \), these vector equations of motion when applied directly do not have the constraint elimination feature of the Lagrange Eq. (20). The objective of Kane and Wang is to modify Eqs. (23) and (24) so as to secure this advantage without accepting the noted limitations inherent in Lagrange's equation in the form of Eq. (20).

To apply the Kane-Wang method, one must first generate the necessary kinematic quantities and record the inertial forces \( \mathbf{F}^{i \ast} \) and inertial torques \( \mathbf{T}^{i \ast} \) for all bodies of the system. This process includes the derivation of the inertial linear and angular velocities \( \mathbf{v}^i \) and \( \omega^i \) of all bodies of the system, and their expression in terms of the constrained generalized coordinates \( q_1, \ldots, q_N \) and their first derivatives. It is always possible to use the \( \nu \) constraint equations given by Eq. (22) to write all velocities \( \mathbf{v}^i \) and \( \omega^i \) in terms of \( N - \nu \) (say, the first \( N - \nu \) generalized velocities \( \dot{q}_{1}, \ldots, \dot{q}_{N-\nu} \)), retaining in general all \( N \) coordinates \( q_1, \ldots, q_N \) in these expressions.

The next step is the selection (by inspection of the expressions for \( \mathbf{v}^i \) and \( \omega^i \)) of \( N - \nu \) variables \( u_1, \ldots, u_{N-\nu} \) so that (1) each \( u_i \) may be written as a linear combination of the \( N - \nu \) generalized velocities \( \dot{q}_{1}, \ldots, \dot{q}_{N-\nu} \) plus a residual term free of generalized velocities, i.e.,

\[
u_i = \sum_{j=1}^{N-\nu} U_{ij} \dot{q}_j + U_{i,} \quad i = 1, \ldots, N - \nu
\]

(27)

(where \( U_{ij} \) and \( U_i \) depend on \( q_1, \ldots, q_N \) and \( t \), and \( \dot{q}_j \) each velocity \( \mathbf{v}^i \) and angular velocity \( \omega^i \) may similarly be written as a linear combination of the variables \( u_1, \ldots, u_{N-\nu} \), i.e.,

\[
\mathbf{v}^i = \sum_{j=1}^{N-\nu} v_{ij} u_j + v_i^i
\]

(28)

\[
\omega^i = \sum_{j=1}^{N-\nu} \omega_{ij} u_j + \omega_i^i
\]

(29)

This step clearly involves some judgment on the part of the analyst, but in specific applications the choice of variables \( u_i \) is not difficult. It is always possible to choose \( u_i = \dot{q}_i, \) \( i = 1, \ldots, N - \nu \), but this is rarely the most attractive choice. More often a judicious choice of angular velocity measure numbers for \( u_1, \ldots, u_{N-\nu} \) is useful. Examples in Refs. 26 and 27 are helpful in establishing a rationale for this selection.
From this point, the application of the method is quite routine. The vectors $v_i^j, \omega_i^j, v_i^{k+1}, \cdots, v_i^{n}, \omega_i^k, \cdots, \omega_i^{n}$ $(i = 1, \cdots, n)$ are defined by the selection of variables $u_i, \cdots, u_{n-x}$ in Eqs. (28) and (29). These vectors are now used to define the "generalized active forces"

$$K_j = \sum_{i=1}^{n} (v_i^j \cdot F^i + \omega_i^j \cdot T)^i, \quad j = 1, \cdots, N - \nu$$

(30)

and the "generalized inertia forces"

$$K_j^* = \sum_{i=1}^{n} (v_i^j \cdot F^{i*} + \omega_i^j \cdot T^{i*}), \quad j = 1, \cdots, N - \nu$$

(31)

Kane's theorem (proven in Ref. 26) has the form

$$K_j + K_j^* = 0, \quad j = 1, \cdots, N - \nu$$

(32)

Kane shows that the basic first-order dynamic equation (32) does not involve the unknown forces and torques required to maintain the constraints of Eq. (22). Thus the combination of the $N - \nu$ dynamic equations of Eq. (32), the $\nu$ constraint equations of Eq. (22), and the $N - \nu$ kinematic equations of Eq. (27) constitutes a complete set of $2N - \nu$ first-order differential equations. This is the minimum number of equations consistent with the selection of a system of $N$ generalized coordinates subject to $\nu$ constraints. In application to the point-connected sets of $n$ rigid bodies of Section II-B, $N$ is $3n$, so Kane's equations are the same in number as those obtained by Russell's approach (Ref. 19), and the same as those actually integrated in Fleischer's program (Ref. 18). If the constraint equations are holonomic (so Eq. 22 is integrable), it is possible to replace $\nu$ of the first-order differential equations by algebraic equations. When these can be solved explicitly for $\nu$ coordinates in terms of the $N - \nu$ remaining coordinates, the results can be substituted into the dynamic equation (32) and the kinematic equation (27) to obtain a minimum set of $2(N - \nu)$ first-order differential equations.

D. Summary

The objective here has been to outline the several approaches to the formulation of equations of motion of discrete-parameter systems in sufficient detail to develop some appreciation of the differences in points of view, and to stimulate interest in deeper inquiry. For a comprehensive treatment of the subject, the reader is referred to the cited works.

It is impossible to make a definitive judgment of the relative merits of the many available approaches. It is difficult to weigh computer time against analyst time, particularly in the face of absolute time constraints and limitations of personnel assigned to a given task at a given time. Even with a given "cost function," the optimum approach varies from one dynamic system to the next.

Within the framework of fully programmed multipurpose digital computer programs, which require the minimum of analyst time and thought, the work of Fleischer (Ref. 18) is perhaps most useful, since it combines many of the attractive features of the earlier works of Hooker and Margulies (Ref. 14), Roberson and Wittenberg (Refs. 15, 16), and Velman (Ref. 20). It is possible that Fleischer's program would run faster if relative motion coordinates were employed, but this would introduce the judgment of the analyst into the simulation process.

There may be some question concerning the desirability in some organizations of developing a multipurpose program. Russell (Ref. 19) and Kane and Wang (Refs. 26 and 27) offer procedures that seem to lead to more efficient simulations than are available with the more highly formalized methods cited above, but they require individual programming of each problem by a capable analyst, which may be an unacceptable constraint.

III. Hybrid-Coordinate Method

A. Vehicle Mathematical Model

The concept of the hybrid-coordinate method depends upon the possibility of separating a given vehicle into a number of idealized structural subsystems, each of which may be classified either as a flexible appendage or as a rigid body or particle. A flexible appendage is by present definition a linearly elastic structure (developing elastic stresses in proportion to strains) for which "small" deformations may be anticipated (so elastic stresses remain proportional to deformations). In most cases, internal energy dissipation may be represented by modal damping (Ref. 30), although discrete dampers can be accommodated. The usual definitions of rigid body and particle are retained.

In its present stage of development, this method is formally restricted by the requirement that each flexible appendage be attached only to a rigid body, or to more than one rigid body if those bodies are themselves interconnected in such a way that their relative motion cannot induce deformation of the flexible appendage. Thus the
appendage cannot properly be the only connecting structure between two rigid bodies, but it can be hinged to two bodies if their relative motion is restricted so as to permit the appendage to accommodate their relative motion by rotating without deforming. Figure 3 illustrates several examples of idealized mechanical assemblages that might present real or imagined obstacles in modeling for hybrid-coordinate analysis. In each of these sketches the structure drawn as a truss or gridwork is supposed to be substantially more flexible than the solid components.

Figure 3a bears some resemblance to a dual-spin vehicle with a flexible antenna, two flexible solar-cell arrays, and two flexible booms on the rotor. Here the de-spun platform $B_1$, the nutation damper mass $B_2$, the rotor $B_3$, and the contents of a spherical tank of fluid $B_4$ are idealized as rigid. Substructures $A^1, \cdots, A^5$ are idealized as elastic appendages. An arbitrary control system determines the relative rotation of $B_1$ and $B_2$, with the use of data from sensors anywhere on the vehicle. In addition, the entire antenna structure $A^1$ may be controlled to rotate on the base $B_1$ without exceeding the capacity of the method. But if the dish portion of $A^1$ is controlled to rotate relative to its support tower, the model must be modified if the equations to follow are to be applied. An individual appendage is permitted only elastic deformations, so antenna dish and tower could not together be one appendage if large relative rotations are prescribed. Yet they cannot be treated as two separate flexible appendages, since each appendage must be attached to a rigid body (and not to another appendage). The only recourse is to model the tower as one or more rigid bodies, in which case the derivations of the following sections continue to apply.

Similarly, in Fig. 3b one must treat the two flexible antenna structures, together with their interconnecting rigid body, as one flexible appendage (as indicated by the dashed lines). A given flexible appendage cannot be attached to two rigid bodies for which relative motion deforms the appendage, so the massive block between the two antenna dishes cannot be isolated as a rigid body, and must instead be absorbed as part of the larger appendage $A$.

Again in Fig. 3c one is strictly precluded from treating each of the three blocks in the system as a rigid body; the block shown within the dashed lines must be included as a part of the appendage. Because of the large relative motions permitted between $B_1$ and $B_2$, one does not have the option of including $B_1$ plus $B_2$ with the flexible structure instead of the block indicated. If an articulated element such as $B_2$ were attached as well to the block shown within the dashed lines, the hybrid-coordinate method as developed in this report would be applicable only by ignoring the masses on one end of the appendage and, instead, applying appropriate forces and torques to the end. This represents some distortion of the method, and jeopardizes the rationale to be applied to modal-coordinate truncation.

Figure 3d illustrates one of the ways in which a flexible appendage can be attached to two rigid bodies. Constraints between bodies $B_1$ and $B_2$ are such that their relative motion results only in the rotation (not in the deformation) of the appendage $A$. It would not be desirable in this case simply to include the little body $B_2$ as part of the flexible appendage because of the discrete damper connecting $B_2$ to $B_1$. The attachment between a flexible appendage and a rigid body must be consistent with the assumption of modal damping of the appendage if modal coordinates are to be obtained for second-order equations.

When two different points of a flexible structure are attached by discrete dampers (e.g., dashpots) to a rather rigid base (as in Fig. 3e), one might assume that an additional rigid body (such as $B_3$ in Fig. 3e) connects the support points. If this assumption is unacceptable, and a discrete damping mechanism must be incorporated within the appendage, then one can find modal coordinates only after rewriting the equations of motion as first-order equations. Although modal transformations of first-order equations with arbitrary damping are treated in Section III-D, it is tentatively assumed in Section III-B that any damping in the appendage may be modeled as classic modal damping (Ref. 30).

It is evident that the equations in this section will not apply to completely general models of space vehicles, and even when applicable they may not constitute the optimum approach to simulation. For the system of Fig. 3c, for example, it may be preferable to ignore entirely the mass in the truss structure and to treat the vehicle as three discrete rigid bodies, perhaps even in a point-connected topological tree configuration if the truss is short and longitudinally stiff. A discrete-parameter simulation of the vehicle in Fig. 3a would, on the other hand, be hopelessly inefficient, and the results would be inferior to a much less expensive hybrid-coordinate simulation.
Fig. 3. Examples of models for hybrid-coordinate analysis
the vehicles of Fig. 3, although for restricted modes of motion one could obtain vehicle normal coordinates for the corresponding first-order state equations. This procedure might, for example, prove optimum for restricted motions of the mechanisms shown in Figs. 3d and 3e.

In reading the present section it may be helpful to keep Fig. 3a in mind, since this is the kind of system for which the hybrid-coordinate approach is most ideally suited.

In what follows, attention is focused first upon an individual flexible appendage A attached to a single base B. In the exceptional case (e.g., Fig. 3d), where a second rigid body is attached to the appendage, the influence of this body is felt in changing the attitude of the primary base and the appendage in its undeformed state. This attitude is reflected in a direction cosine matrix C, which is considered time-dependent in the derivations.

After the individual appendage equations are examined in detail, the equations of motion of the balance of the vehicle are considered. There is no attempt to include in this report any general procedure for approaching the derivation of these equations. A class of vehicle somewhat less general than that illustrated in Fig. 3a is treated here in detail, although for more complex systems there may be some advantage in combining some of the discrete-parameter methods of Section II with the appendage equations derived explicitly here.

B. Flexible-Appendage Equations

Consider at the outset the equations of motion of a linearly elastic structure A attached to a base B, assuming that the structure undergoes only "small" deformations while the base motion is arbitrary. ("Small" deformations means in a strict mathematical sense arbitrarily small or vanishing deformations, since terms above the first degree in scalar measures of deformation and deformation time derivatives are ignored completely. In engineering practice, however, this requirement is interpreted quite liberally, and a beam that suffers a tip deflection of 10% of its length is commonly described as having "small" deformations.)

For convenience in derivation and for compatibility with engineering practice, the flexible appendage A is idealized initially as a collection of elastically interconnected, discrete rigid subbodies $A_1, \ldots, A_n$. Damping mechanisms are excluded from this idealization,* but modal damping is incorporated in the equations at a later stage of the derivation, with the introduction of modal coordinates for the appendage. At this point, the equations will lose their apparent restriction to discretely modeled appendages, and will be equally applicable to continuously modeled elastic appendages. In special applications to taut strings and membranes or to uniform beams, plates, or shells, the analyst may find a continuous model most convenient. In the vast majority of space applications, however, a discrete-parameter model is necessary. Frequently the model consists simply of elastically interconnected particles, but the present assumption of interconnected rigid bodies is more general and often more convenient.

Let the set of dextral orthogonal unit vectors $b_1, b_2, b_3$ be fixed in the base B, and the similar set $a_1, a_2, a_3$ be fixed in the reference established by A prior to deformation (i.e., fixed in that portion of A contiguous to rigid body B). When A is elastically connected to B, these sets of vectors may be identical, and at most are related by a constant transformation matrix. But when A can rotate relative to B (as in Fig. 3a), the antenna $A^T$ might rotate relative to the base $B_j$, the transformation matrix relating these vector bases will vary with time according to an independently specified control law. If the unit vectors are written as vector arrays

$$
(a) = \begin{pmatrix} a_1 \\ a_2 \\ a_3 \end{pmatrix}, \quad (b) = \begin{pmatrix} b_1 \\ b_2 \\ b_3 \end{pmatrix}
$$

these arrays may be treated like column matrices in recording the relationship between (a) and (b) in items of the direction cosine matrix $C$, i.e.,

$$
(a) = C (b)
$$

In application it is often convenient to be able to select the basis (a) for each appendage individually, and to select the basis (b) independently (perhaps guided by the principal axes of inertia of the total vehicle, or by

*Terms to be added to the equations of this subsection to accommodate idealized discrete dampers are easily generated, and are accommodated in the coordinate transformations of subsequent subsections.

In this report matrices are not identified by brackets [ ] unless the elements of the matrix are recorded in detail. Vector arrays are always enclosed in braces { }.
design coordinates of a drawing, or by attitude-control axes). Thus even when A does not rotate relative to B, the matrix C may not be the identity matrix E. When A does rotate relative to B, so that C varies with time, it will become necessary either to augment the equations in this section by added dynamic equations or to assume that perfect control results in known time behavior of C.

In devising the discrete-parameter model of A, it is convenient to select basis \{a\} and the individual mass-center principal axis bases of \(A_1, \ldots, A_n\) to be identical.

The first step is the formulation of the equations of motion of a typical body \(A_s\) in form convenient for combination with corresponding equations for all \(n\) bodies. Consequently the vector- dyadic equations of translation and rotation will be written as matrix equations in basis \{a\}.

Let \(P_s\) be the mass center of body \(A_s\), and let \(O'\) be a point fixed in inertial space (see Fig. 4). Then Newton’s second law provides

\[
F^* = m_s \ddot{P}_s
\]

where \(F^*\) is the resultant force applied to \(A_s\), \(m_s\) is the mass of \(A_s\), \(P_s\) is the vector from \(O'\) to \(P_s\), and (as noted previously) each dot over a vector denotes time differentiation in an inertial frame of reference.

As shown in Fig. 4, the inertial position vector \(P_s\) may be written as

\[
P_s = X + c + R + r^s + u^s
\]

where \(X\) is the inertial position vector of the vehicle mass center CM; \(c\) is the vector from CM to the point \(O\) fixed relative to body \(B\) and coincident with CM when the vehicle is in some nominal undeformed configuration; \(R\) is the vector from \(O\) to an arbitrary point \(Q\) fixed in \(B\) on the interface between \(A\) and \(B\); \(r^s\) is the vector locating from \(Q\) the point \(Q^s\) occupied by \(P_s\) when \(A\) is undeformed; and \(u^s\) defines the translational deformation of the appendage at point \(Q_s\). The vectors in Eq. (36) are not all expressed conveniently in anyone vector basis. Vector \(X\) defines the vehicle trajectory, which may be known in terms of an inertial reference; \(R\) and \(r^s\), on the other hand, are fixed in the reference frames in which \(b_1, b_2, b_3\) and \(a_1, a_2, a_3\) are embedded, respectively, so they are most efficiently expressed in their own natural vector bases.

Since a matrix formulation is ultimately required, 3 by 1 matrices are defined for each of the vectors in Eq. (36) in terms of the appropriate vector basis. These matrices are introduced with the use of vector arrays (as in Eq. 33), by the definitions

\[
X = \begin{bmatrix} X_1 \\ X_2 \\ X_3 \end{bmatrix} = \begin{bmatrix} i_1 \\ i_2 \\ i_3 \end{bmatrix} X
\]

\[
c = \begin{bmatrix} c_1 \\ c_2 \\ c_3 \end{bmatrix} = \begin{bmatrix} b_1 \\ b_2 \\ b_3 \end{bmatrix} c
\]

\[
R = \begin{bmatrix} R_1 \\ R_2 \\ R_3 \end{bmatrix} = \begin{bmatrix} b_1 \\ b_2 \\ b_3 \end{bmatrix} R
\]

\[
r^s = \begin{bmatrix} r^s_1 \\ r^s_2 \\ r^s_3 \end{bmatrix} = \begin{bmatrix} a_1 \\ a_2 \\ a_3 \end{bmatrix} r^s
\]

\[
u^s = \begin{bmatrix} u^s_1 \\ u^s_2 \\ u^s_3 \end{bmatrix} = \begin{bmatrix} a_1 \\ a_2 \\ a_3 \end{bmatrix} u^s
\]

where the superscript \(T\) denotes the transpose of a matrix or vector array.

Upon substituting Eq. (36) into Eq. (35), one confronts the necessity of differentiating in an inertial reference frame a number of vectors that are much more easily expressed (and sometimes constant) in some other reference frame. This is most easily accomplished by liberal use of the identity from vector differential calculus

\[
\frac{d}{dt} V = \frac{d}{dt} V + \frac{d}{dt} \omega f_s \times V
\]

where \(V\) is any vector, \(\frac{d}{dt} \omega f_s\) is the angular velocity of any reference frame \(f_s\) relative to any other reference frame \(f_t\), and the superscript preceding the derivative operator denotes the reference frame of differentiation. Although \(t\) may be any independent variable, in the present context it is always time.
If the presuperscript \( i \) denotes the inertial reference frame, Eqs. (35) and (36) may be combined as

\[
F^i = m_i \left[ \frac{d^2}{dt^2} X + \frac{d^2}{dt^2} (c + R) + \frac{d^2}{dt^2} (r^i + u^i) \right]
\]  
(39)

The first term in brackets in Eq. (39) is (with the use of Eq. 37) simply

\[
\frac{d^2}{dt^2} ((i)^r X) = (i)^r \ddot{X}
\]  
(40)

since the inertial time derivative of \( (i) \) is zero. (Recall that a matrix derivative such as \( \ddot{X} \) is just the matrix of differentiated elements.)

The second term in brackets in Eq. (39) may be obtained (with the use of Eq. 38) from

\[
\frac{d}{dt} (c + R) = \frac{d}{dt} (c + R) + \omega \times (c + R)
\]  
(41)

where \( b \) is the reference frame of body \( B \) and basis \( \{b\} \), and \( \omega \) is the inertial angular velocity of \( b \). Differentiating Eq. (41) provides (with the use of Eq. 38 again)

\[
\frac{d^2}{dt^2} (c + R) = \frac{d}{dt} \left( \frac{d}{dt} (c + R) + \omega \times (c + R) \right)
\]  
(42)

Now Eq. (37) may be utilized, noting \( R \) to be constant, to obtain

\[
\frac{d^2}{dt^2} (c + R) = (b)^r \ddot{c} + 2\omega \times (b)^r \dot{c} + \omega \times [\omega \times (b)^r (c + R)] + \dot{\omega} \times (b)^r (c + R)
\]  
(43)

To express this result in matrix terms, one must obtain the matrix equivalent of a vector cross product.

For two arbitrary vectors \( V \) and \( W \), expressed in terms of an arbitrary vector basis \( \{e\} \), the vector cross product may be written as

\[
V \times W = (e)^r V \times (e)^r W = (e)^r \tilde{V} W
\]

where

\[
V = \begin{bmatrix} V_1 \\ V_2 \\ V_3 \end{bmatrix}, \quad W = \begin{bmatrix} W_1 \\ W_2 \\ W_3 \end{bmatrix},
\]

\[
\tilde{V} = \begin{bmatrix} 0 & -V_3 & V_2 \\ V_3 & 0 & -V_1 \\ -V_2 & V_1 & 0 \end{bmatrix}
\]  
(44)

as may be confirmed by expansion. In equivalent terms, the matrix product \( \tilde{V} W \) is isomorphic to the vector product \( V \times W \). Thus the matrix identity

\[
\tilde{V} W = -\tilde{W} V
\]  
(45)

follows from the vector identity \( V \times W = -W \times V \). The tilde operator \( (\sim) \) over a 3 by 1 matrix represents the
corresponding skew-symmetric 3 by 3 matrix, as in Eq. (44), throughout this report. Note the useful identity
\[ \vec{V} = -\vec{V} \]  
(46)
for any tilded matrix, because of skew symmetry.

Equation (43) can now be written (with the definition \[ \omega \equiv (b) r_\omega \]) as
\[ \frac{d^2}{dt^2} (r^* + u^*) = \frac{d^2}{dt^2} u^* + 2\Omega^2 \times \frac{d}{dt} u^* 
+ \Omega_\omega \times [\Omega_\omega \times (r^* + u^*)] 
+ \frac{b d}{dt} \Omega^2 \times (r^* + u^*) 
+ 2\omega \left( \frac{d}{dt} u^* + \Omega \times (r^* + u^*) \right) 
+ \omega \times [\omega \times (r^* + u^*)] + \dot{\omega} \times (r^* + u^*) \] 
(48)
where \( \Omega^2 \) is the angular velocity of the \( (a) \) frame relative to the \( (b) \) frame, and it has been recognized that the time derivative of \( r^* \) in reference \( a \) is zero. This result is now to be written in terms of the vector array \( (a) \) and the matrices defined in Eq. (37). If \( \Omega^2 \) is written in the \( (a) \) basis as
\[ \Omega^2 = (a)^r \Omega^2 \] 
(49)
and Eq. (34) is used to write
\[ (b)^r = (a)^r C_\omega \] 
(50)
so that \( \omega \) is written in the \( (a) \) basis as
\[ \omega = (b)^r \omega = (a)^r C_\omega \] 

The result of these substitutions into Eq. (39) is the vector equation
\[ (a)^r F^* = (a)^r m_s \left( C_\omega \vec{X} + C_\omega [\vec{c} + 2\vec{c} \dot{c} + \vec{c} (c + R) + \vec{c} (c + R)] + \dot{u}^* + 2\tilde{u}^* \tilde{u}^* + \tilde{u}^* \tilde{u}^* (r^* + u^*) 
+ \tilde{u}^* (r^* + u^*) + 2 (C_\omega) \tilde{u}^* + 2 (C_\omega) \tilde{u}^* (r^* + u^*) + (\tilde{C}_\omega) (\tilde{C}_\omega) (r^* + u^*) + (\tilde{C}_\omega) (\tilde{C}_\omega) (r^* + u^*) \right) \] 
(54)
When Eqs. (51), (47), and (40) are combined in Eq. (39), the three different vector bases of these equations are abandoned in favor of basis \( (a) \), noting that this is also the natural basis for \( F^* \). This requires the use of Eq. (50) in Eq. (47), and the parallel use in Eq. (40) of the relationship
\[ (i)^r = (b)^r \Theta = (a)^r C_\Theta \] 
(52)
where \( \Theta \) is the direction cosine matrix relating the body \( B \) to an inertially fixed vector basis, i.e.,
\[ (b) = \Theta (i) \] 
(53)

The result of these substitutions into Eq. (39) is the vector equation
\[ (a)^r F^* = (a)^r m_s \left( C_\Theta \vec{X} + C_\Theta [\vec{c} + 2\vec{c} \dot{c} + \vec{c} (c + R) + \vec{c} (c + R)] + \dot{u}^* + 2\tilde{u}^* \tilde{u}^* + \tilde{u}^* \tilde{u}^* (r^* + u^*) 
+ \tilde{u}^* (r^* + u^*) + 2 (C_\Theta) \tilde{u}^* + 2 (C_\Theta) \tilde{u}^* (r^* + u^*) + (\tilde{C}_\Theta) (\tilde{C}_\Theta) (r^* + u^*) + (\tilde{C}_\Theta) (\tilde{C}_\Theta) (r^* + u^*) \right) \] 
(54)
which employs the new matrix symbol $F'$ as defined by

$$F' = \{a\}^T F$$  \hspace{1cm} (55)$$

One may now record the corresponding matrix equation (in vector basis $\{a\}$), either by inspection of Eq. (54), or by formal dot-multiplication by the vector array $\{a\}$, noting that the relationship

$$\{a\} \cdot \{a\}' = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} = E$$

follows from the introduction of the dot between vector arrays to imply the multiplication of vector arrays following the rules of matrix multiplication, but with dot-multiplication of the vector elements inside the arrays. This notation is sometimes convenient for notational bookkeeping, but even without this artifice one can simply record the matrix equation isomorphic to the vector equation (54). In the process, the future direction of the derivation may be anticipated by the replacement of the scalar $m_\iota$ with the matrix $m' = m_\iota E$, with $E$ the 3 by 3 identity matrix. The desired matrix equation is then obtained (after some factoring and rearrangement) in the form

$$F' = m' \left\{ C [\hat{\phi} \dot{X} + \ddot{c} + 2 \hat{\phi} c + \hat{\phi} \omega (c + R) - (\hat{\phi} + \tilde{\phi}) \ddot{\omega} - C \dot{\omega} \right\} + \left[ (\tilde{C}_\omega \tilde{C}_\omega^T + 2 (\tilde{C}_\omega \tilde{C}_\omega^T \ddot{\omega}) + \tilde{C}_\omega \tilde{C}_\omega^T \dddot{\omega} \right] \right\} (r' + u') + 2 \left[ \tilde{C}_\omega + (\tilde{C}_\omega^T) \right] \dot{u}' + \ddot{u}'$$

\hspace{1cm} (57)

Of course the matrix $c$, which denotes the translation in body $B$ of the vehicle mass center, depends in part on the translations $u^s (s = 1, \ldots, n)$ of the sub-bodies of the appendage $A$. In general, it may also depend on the deformations of other elastic appendages of the system, and on the (possibly large) relative motions of other articulated rigid bodies of the vehicle. To emphasize the influence on $c$ of the deformation of the appendage for which equations of motion are being written, the mass-center definition is employed to write for $c$

$$c = -\frac{1}{\Omega n} \sum_{s=1}^{n} m_s u^s + e$$

\hspace{1cm} (58)

where $e$ accommodates the mass-center motion due to other appendages and moving parts, if any exist, and $\Omega n$ is the total vehicle mass. With the definition

$$e = \{b\}^T e$$

\hspace{1cm} (59)

Eq. (58) may be written in the form

$$\{b\}^T c = -\frac{1}{\Omega n} \sum_{s=1}^{n} \left\{ \sum_{s=1}^{n} m_s u^s \right\} (a)^T e$$

$$= \{b\}^T \left[ -\sum_{s=1}^{n} C_s^r \mu^s u^s + e \right]$$

where $\mu^s = m^s / \Omega n$ and Eq. (34) has been employed. This vector equation has the matrix counterpart

$$c = -\sum_{s=1}^{n} C_s^r \mu^s u^s + e$$

\hspace{1cm} (60)

Thus the role of the appendage deformation in Eq. (57) is made more explicit by the substitution of Eq. (60) and its derivatives (permitting $C$ to vary with time), which results in

$$\dot{c} = -\sum_{s=1}^{n} (C_s^r \mu^s u^s + C_s^r \mu^s \dot{u}^s) + \ddot{c}$$

\hspace{1cm} (61)

and

$$\dddot{c} = -\sum_{s=1}^{n} (C_s^r \mu^s u^s + 2 C_s^r \mu^s \dot{u}^s + C_s^r \mu^s \ddot{u}^s) + \dddot{c}$$

\hspace{1cm} (62)

If it seems preferable to avoid the time derivatives of the direction cosine matrix $C$, this can be accomplished by using the identity (38) once more to obtain

$$\{b\}^T \dot{c} = \frac{b}{dt} \left[ (\{b\}^T c) \right]$$

$$= \frac{b}{dt} \left[ \left\{ \sum_{s=1}^{n} m_s u^s \right\} (a)^T e \right]$$

\hspace{1cm} (63)

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With Eq. (34), this yields the matrix equation

$$
\dot{\mathbf{c}} = -\sum_{s=1}^{n} C^T \mu^s \dot{u}^s - C^T \Omega^s \sum_{s=1}^{n} \mu^s u^s + \dot{\mathbf{c}}
$$

(64)

Differentiation of Eq. (63) in reference frame \( b \) similarly provides the vector equation

$$(\mathbf{b})^T \dot{\mathbf{c}} = - (\mathbf{a})^T [ \sum_{s=1}^{n} \mu^s \dot{u}^s + \Omega^s \sum_{s=1}^{n} \mu^s u^s ] - (\mathbf{a})^T \Omega^s [ \sum_{s=1}^{n} \mu^s \dot{u}^s + \Omega^s \sum_{s=1}^{n} \mu^s u^s ] + (\mathbf{b})^T \dot{\mathbf{c}}
$$

The matrix counterpart in basis \( \{b\} \) for this vector equation is

$$
\dot{\mathbf{c}} = - C^T [ \sum_{s=1}^{n} \mu^s \dot{u}^s + \Omega^s \sum_{s=1}^{n} \mu^s u^s + 2 \Omega^s \sum_{s=1}^{n} \mu^s \dot{u}^s + \Omega^s \Omega^s \sum_{s=1}^{n} \mu^s u^s ] + \dot{\mathbf{c}}
$$

(65)

Now Eqs. (64) and (65) may be used in Eq. (57), instead of substituting Eqs. (61) and (62) in this equation, thereby avoiding derivatives of \( C \). Alternatively, one can avoid the relative angular rate \( \Omega^s \) and its time derivatives entirely by the substitution of the kinematic identity (Ref. 16)

$$
\Omega^s = C \dot{C}^T
$$

(66)

and its consequence

$$
\dot{\Omega}^s = \dot{C} \dot{C}^T + C \dot{C}^T
$$

(67)

With these substitutions in Eq. (57), and with Eqs. (61) and (62), all relative motion between bases \( \{a\} \) and \( \{b\} \) is expressed in terms of the direction cosine matrix \( C \) and its derivatives. Although this option may be optimum in some applications, it would seem preferable in most practical cases to avoid \( \dot{C} \) and \( \dot{C}^T \), and therefore to employ Eqs. (57), (64), and (65).

Since the sub-bodies \( A_1, \ldots, A_n \) are considered to be rigid bodies rather than particles, the translational equation (67) must be augmented by rotational equations such as

$$
\mathbf{T'} = \dot{\mathbf{H}}^s
$$

(68)

for typical rigid sub-body \( A_s \). Here \( \mathbf{H}^s \) is the inertial angular momentum of \( A_s \) referred to the mass center \( P_s \), and \( \mathbf{T'} \) is the corresponding resultant torque. The rules of vector differentiation (see Eq. (38)) may now be applied to \( \mathbf{H}^s \), or to its equivalent \( \mathbf{I'} \cdot \omega^s \), where \( \mathbf{I'} \) is the mass-center inertia dyadic of \( A_s \) and \( \omega^s \) the inertial angular velocity of \( A_s \). The result is

$$
\frac{\mathbf{I'}d}{dt} \mathbf{D} = \frac{\mathbf{I'}d}{dt} \mathbf{D} + \mathbf{I'} \omega^s \times \mathbf{D} - \mathbf{D} \times \mathbf{I'} \omega^s
$$

(70)

with \( \mathbf{D} \) any dyadic and \( f_1 \) and \( f_2 \) any two frames of reference. (This relationship can be confirmed by writing the dyadic in the form \( \mathbf{D} = \mathbf{D}_{ef} \mathbf{e}_s \mathbf{e}_p \) and then applying Eq. (38) to the unit vectors \( \mathbf{e}_a \) and \( \mathbf{e}_p \).)

The term \( \mathbf{I'} \times \mathbf{I'} \omega^s \times \omega^s \) in Eq. (69) is zero, since the implied operations include dot-multiplying by \( \omega^s \) a vector orthogonal to \( \omega^s \). Equation (69) is further simplified by the constancy of the inertia dyadic of the rigid body \( A_s \) in the reference frame of that body, i.e., \( \mathbf{I'} \omega^s \times \mathbf{I'} \) is zero. In rewriting the correspondingly simplified rotational equation, one may expand the inertial angular velocity \( \omega^s \) of \( A_s \) by use of the "chain rule"

$$
\mathbf{I'} \omega^s = \mathbf{I'} \omega^s + \mathbf{I'} \omega^s + \cdots + \mathbf{I'} \omega^s
$$

(71)

*Throughout this report, lower-case Greek indices range in value from 1 to 3, and when these indices are repeated in a given term, summation over these values is implied.*
where $f_1, \ldots, f_n$ are $n$ arbitrary frames of reference and $f^\omega$ is the angular velocity of frame $f_i$ relative to frame $f_j$. In this case the chain is from the frame of body $A_s$ to the frame of $A$, to the frame of $B$, to inertial space. Since the rotation of $A_s$ relative to $A$ is due only to "small" structural deformations, this rotation can be represented by the vector $\beta^s = \beta_1^s a_1 + \beta_2^s a_2 + \beta_3^s a_3$, where $\beta_1^s, \beta_2^s, \beta_3^s$ are angles of rotation about orthogonal axes $a_1, a_2, a_3$. This representation is strictly applicable only if these rotations are infinitesimal, but this is nonetheless the commonplace assumption of structural dynamicists. Consequently $w$ in Eq. (69) can be replaced by the use of Eq. (71) to write

$$w = \omega + \Omega e + (a)^T \beta^s$$

If all vectors are now written in terms of the vector arrays of the most convenient basis, with the use of the definitions following Eq. (49) together with

$$T^* = (a)^T T^*$$

and, with the summation convention, and new unit vectors $a^*_1, a^*_2, a^*_3$ embedded in $A_s$ as the principal axis vector basis

$$T^* = I^* a^*_1 a^*_2 a^*_3 = (a^*_1 a^*_2 a^*_3)\begin{bmatrix} I_{11} & 0 & 0 \\ 0 & I_{22} & 0 \\ 0 & 0 & I_{33} \end{bmatrix} (a^*_1)$$

the vector rotational equation becomes

$$(a)^T T^* = (a)^T (a) \cdot [(b)^T \Omega + (a)^T \Omega e + (a)^T (a)^T \beta^s + ((b)^T \Omega + (a)^T \Omega e) \times (a)^T \beta^s]$$

$$+ ((b)^T \Omega + (a)^T \Omega e) \times (a)^T I^* (a)^* \cdot ((b)^T e + (a)^T \Omega e)$$

$$+ ((b)^T e + (a)^T \Omega e) \times (a)^T I^* (a)^* \cdot (a)^T \beta^s \times (a)^T I^* (a)^* \cdot ((b)^T \Omega + (a)^T \Omega e)$$

By using Eq. (50) to obtain every vector in basis $(a)$, and applying the matrix representations of vector cross-multiplication (see Eq. 44) and vector array dot-multiplication (Eq. 56), one can obtain this vector equation in the useful form [noting $(a)^* \equiv (E - \beta^s) (a)$, so $(a)^T = (a)^T (E + \beta^s)$]

$$(a)^T T^* = (a)^T (I^* [C_\omega + \hat{\Theta} e + (\tilde{C}_\omega) \Omega e + \hat{\Theta}^s + (\tilde{C}_\omega) \beta^s + \hat{\Theta}^s \beta^s]) + [(\tilde{C}_\omega) + \hat{\Theta} e] [\hat{\Theta}^s I^* - I^* \hat{\Theta}^s] [(\tilde{C}_\omega) + \hat{\Theta} e] + [(\tilde{C}_\omega) + \hat{\Theta} e] I^* [C_\omega + \Omega e] + [(\tilde{C}_\omega) + \hat{\Theta} e] I^* \beta^s + \hat{\Theta} \beta^s I^* [C_\omega + \Omega e]$$

Now the isomorphic matrix equation can be written by inspection (or obtained formally by dot-multiplying by $(a)$). In the process, the identity of Eq. (45) is applied whenever it seems convenient to have the most obviously unknown variable available for factoring on the extreme right. For example, in the last term the unknown deformation $\beta^s$ must appear on the right if it is to be factored out with the term preceding it, so the identity

$$\hat{\Theta} \beta^s I^* (C_\omega + \Omega e) = - [I^* (C_\omega + \Omega e)]^{-1} \hat{\Theta}^s \beta^s$$
is used. Here the tilde outside the square brackets is to be construed to apply to the entire 3 by 1 column matrix within those brackets. With this substitution, the matrix equation corresponding to Eq. (77) becomes*

\[
T^* = I^* [C^* \dot{\omega} + \ddot{\theta}^*] + [I^* (\dddot{C^*}) + (\dddot{C^*}) I^* + I^* \dddot{\omega}^* \dot{\theta}^* - (I^* \dddot{C^*} + I^* \Omega^* \omega^*) \dot{\theta}^* + \{ - [I^* (\dddot{C^*} \dot{\omega} - \dddot{\omega}^* C^*) \} \beta^* + I^* (\dddot{C^*} \dot{\omega} - \dddot{\omega}^* C^*) C^* - (C^* \omega^* \Omega^* + \omega^* \Omega^* C^*) I^* (C^* \omega^* + \omega^* \Omega^*) \beta^* - I^* \dddot{\omega}^* C^* + [(\dddot{C^*}) + \dddot{\omega}^*] I^* [(C^* \omega^* + \omega^* \Omega^*)] \\
(79)
\]

Again it should be noted that these rotational equations can, with the use of Eqs. (66) and (67), be written in a form not involving \(\Omega^*\), should this be desirable in a special case.

In most applications, it is convenient to accept Eqs. (79) and (57) as the rotational and translational equations of motion, respectively, of body \(A_n\), with Eqs. (60), (64), and (65) substituted into Eq. (57) to accommodate vehicle mass-center motion relative to body \(B\). The result of this substitution is recorded as follows, with the temporary convention that \(\Sigma\) means the sum over \(k\) ranging from 1 to \(n\):

\[
F^* = m^* \{C_0 \dddot{X} - (\dddot{C^*} + \dddot{\omega}^* C + \omega^* \Omega^*) \omega - \dddot{\omega}^* \dot{\theta}^* + \Sigma_{k} \Sigma_{k} \dddot{\omega}^* + C_0 + 2 [(\dddot{C^*}) (C^*)] \dddot{\omega}^* - 2 C_0 C^* \Sigma_{k} \dot{\theta}^* + 2 C_0 \dddot{e} \nonumber \\
- 2 \dddot{\omega}^* \Sigma_{k} \dot{\theta}^* - C_0 C^* \Sigma_{k} \dot{\theta}^* + [(\dddot{C^*}) (C^*)] \dddot{\omega}^* - \dddot{\omega}^* \Sigma_{k} \dot{\theta}^* - 2 C_0 C^* \Sigma_{k} \dot{\theta}^* + \dddot{\omega}^* \dot{\theta}^* - \dddot{\omega}^* \dot{\theta}^* - \dddot{\omega}^* \dot{\theta}^* + \dddot{\omega}^* \dot{\theta}^* + \dddot{\omega}^* \dot{\theta}^* + [(\dddot{C^*}) (C^*)] + 2 (\dddot{C^*}) (\dddot{\omega}^*) + \dddot{\omega}^* (\dddot{\omega}^*) + \dddot{\omega}^* (\dddot{\omega}^*) (r^* + u^*) \\
(80)
\]

Again the identity \(\dddot{C^*} = \dddot{\omega}^* C^*\) might be substituted, and also the consequence \(\dddot{C^*} = \dddot{\omega}^* C^*\).

Equations (79) and (80) constitute the most general formulation of equations of motion of an appendage sub-body \(A_n\) to be derived in this report. These equations are applicable to a sub-body of an appendage that is undergoing small deformations while rotating in an arbitrary way (as described by \(C\) and \(\Omega^*\)) relative to a base \(B\) subject to any translation and rotation (as described by \(\dot{X}\) for the vehicle \(CM\) and \(\theta\) and \(\omega\) for body \(B\)). Any number of additional appendages or articulated moving parts may also be attached to \(B\); these internal motions will influence \(\theta\), \(\omega\), and \(e\).

In practice, the generality of these equations is rarely required. Only in the exceptional case (e.g., the scanning antenna) is there a nonzero \(\Omega^*\), so \(C\) is usually constant. Then there is no theoretical objection to replacing \(C\) by the identity matrix \(E\), particularly if there is but one appendage attached to body \(B\). For reasons not yet apparent, it may be computationally desirable to adopt the view that all bodies \(B\) have only one appendage, even though the appendage may in some cases be composed of several physically distinct structures attached to \(B\). Then \(e\) is zero, except when a statically unbalanced rotating rigid body is also attached to \(B\), or a secondary rigid body can translate relative to \(B\).

For the simplest configuration, with \(C = E\) and \(e = 0\), the equations of motion (79) and (80) adopt the form

\[
T^* = I^* [\dot{\omega}^* + \dddot{\theta}^*] + [I^* \dddot{\omega}^* + \dddot{\omega}^* I^* - (\dddot{P^*} \omega)] \dot{\theta}^* + \dddot{\omega}^* I^* \omega \\
+ [I^* \dddot{\omega}^* - (\dddot{P^*} \omega) - \dddot{\omega}^* (\dddot{P^*} \omega) + \dddot{\omega}^* I^* \omega] \dot{\theta}^* \\
(81)
\]

The identity \(\dddot{C^*} = \dddot{\omega}^* C^*\) should perhaps be noted, although this substitution into Eq. (79) seems computationally inefficient.
\[ F^s = m^s \left[ \ddot{\mathbf{X}} + \ddot{\mathbf{u}}^s - \Sigma_{j,k} \ddot{u}_j^k + 2 \dddot{u}^s \right] \\
- \left( \mathbf{R} + \mathbf{P} \right) \dot{\mathbf{u}} + \dddot{u}^s \left( \mathbf{R} \right) \\
+ \ddot{\alpha} \left[ \mathbf{R} \mathbf{r}^s + \mathbf{u}^s - \Sigma_{j,k} \mathbf{u}^k \right] \]  

(82)

These equations still permit unrestricted motion of base B.

The forces \( F^s \) and torques \( T^s \) that appear in the basic differential equations (79) and (80) of sub-body \( A_s \) include terms due to structural interactions with neighboring sub-bodies of the appendage \( A \). These interactions are represented here as linearly elastic and viscous forces and torques, so they are proportional to the deformations and deformation rates as represented by \( u^1, \ldots, u^n, \beta^1, \ldots, \beta^n \) and their first derivatives. In practice, viscous damping terms are often unspecified until transformation to modal coordinates has been accomplished.

The resulting approximation is far from perfect in its representation of structural behavior. Even when the materials of a complex structure are essentially linearly elastic, there may be sufficient "play" in the joints of the assembled structure to jeopardize the assumption of linear elasticity of the composite structure. Furthermore, the assumption of viscous damping is not strictly in conformity with the performance of even the simplest structural elements. But for complex structures lacking discrete artificial dampers, the assumptions of linear elasticity and viscous modal damping are almost universal practice in structural dynamics, and these assumptions are reasonably well supported by vibration test data (Refs. 31, 32). They are accordingly the assumptions adopted in this report.

Structural forces and torques in each of the equations typified by Eqs. (79) and (80) therefore may couple these equations to those of every other sub-body of the appendage. The most convenient method of recording all the coupled translational and rotational equations of the \( n \) bodies \( A_n, \ldots, A_s \) is with a single matrix equation of dimension \( 6n \).

Because interest is focused presently on the appendage deformations \( u^1, \ldots, u^n, \beta^1, \ldots, \beta^n \), only terms in these variables will be written on the left side of the appendage matrix equation. The single \( 6n \) by 1 matrix

\[ q = [u^1, u^2, u^3, \beta^1, \beta^2, \beta^3, \ldots, \beta^n]^T \]  

(83)

fully characterizes the appendage deformations.

Because the equations have been linearized in the appendage deformations, the matrix equation must have the structure

\[ M^' \ddot{q} + D' \dot{q} + G' q + K' q + A' q = L' \]  

(84)

where by definition \( D' \) and \( K' \) are symmetric and \( G' \) and \( A' \) are skew-symmetric. Inspection of Eqs. (79) and (80) reveals that \( M' \) is a constant symmetric matrix, but \( G', A', \) and \( K' \) depend on the variables \( \omega, \Omega, \) and \( C \). The matrix \( L' \) depends on these variables and in addition on \( \Theta, \mathbf{X}, e, \) and the external applied forces that may appear in \( F^* \) and \( T^* \). The matrix \( D' \) accommodates damping in the structure.

The detailed representation of the \( 6n \) by \( 6n \) coefficient matrices \( M', G', A', \) and \( K' \) and the \( 6n \) by 1 matrix \( L' \) is facilitated by the introduction of the Boolean operator matrices

\[ \Sigma_{S0} = \begin{bmatrix} E & 0 & 0 & \cdots & E & 0 \end{bmatrix} \] 

\[ \Sigma_{Sg} = \begin{bmatrix} 0 & 0 & 0 & \cdots & 0 & E \end{bmatrix} \]  

(85)

where \( E \) and 0 are 3 by 3 matrices, the former being the unit matrix and the latter the null matrix. In the present application the matrices \( \Sigma_{S0} \) and \( \Sigma_{Sg} \) will generally have \( 6n \) rows, but this restriction is not embodied in the definition, since the matrix dimension is always clear from the context in which it is used.

The operators \( \Sigma_{S0} \) and \( \Sigma_{Sg} \) serve to distribute a given 3-row matrix into alternate 3-row partitions of a larger matrix, having perhaps \( 6n \) rows. For example, the expression \( \Sigma_{S0} \mathbf{C} \mathbf{X} \) is the column matrix

\[ [(\mathbf{C} \mathbf{X}) 0 (\mathbf{C} \mathbf{X}) 0 \cdots (\mathbf{C} \mathbf{X}) 0]^T \]

which, by inspection of Eqs. (79) and (80), must appear as part of \( L' \) in Eq. (84).

The operators \( \Sigma_{S0} \) and \( \Sigma_{Sg} \) are, on the other hand, effectively selective summing operators, since, for example, when multiplied by a \( 6n \)-row matrix they sum respectively the odd- and even-numbered partitioned 3-row submatrices of the \( 6n \)-row matrix into a single 3-row matrix. As a specific example, \( \Sigma_{S0} ^* \hat{q} \) provides the 3 by 1 column matrix obtained by adding \( \ddot{u}^1 + \ddot{u}^2 + \cdots + \ddot{u}^n \).
The constant matrix $M'$ in Eq. (84) is now most easily written in terms of the new symbol $M$, defined in terms of 3 by 3 partitioned matrices as

$$
M = \begin{bmatrix}
    m^1 & 0 & 0 & 0 \\
    0 & I^1 & 0 & 0 \\
    0 & 0 & m^2 & 0 \\
    0 & 0 & 0 & I^2 \\
    \vdots & \vdots & \vdots & \vdots \\
    0 & 0 & 0 & 0 \\
\end{bmatrix}
$$

(86)

Now $M'$ can be recorded as

$$
M' = M (E - \Sigma_{n=0}^N M_n) 
$$

(87)

Here $E$ is an identity matrix of dimension $6n$ by $6n$, and $\Omega$ is, as previously, the total vehicle mass (recall $\mu = m^2/\Omega$). The matrix $M$ may be recognized as the inertia matrix that would be involved in a calculation of cantilever modes of the appendage. This interpretation will be explored in detail in Section III-D on coordinate transformations.

In expansion, $M'$ appears in terms of 3 by 3 partitioned matrices as shown in Fig. 5. Note that $M'$ is a symmetric matrix of constants.
Before recording the matrix $G'$ in expanded form, it is useful to examine the coefficient of $\hat{\beta}^s$ in Eq. (79) in some detail. It is convenient to define the new symbol $\omega_s = \Omega^s + C_\omega$. Then, the coefficient of $\hat{\beta}^s$ becomes

$$
I \tilde{\omega}^s + \tilde{\omega}^s I - (I \tilde{\omega}^s) = \begin{bmatrix}
0 & -(I_1 + I_2 - I_3) \omega_3^s & (I_1 - I_2 + I_3) \omega_3^s \\
(I_1 + I_2 - I_3) \omega_3^s & 0 & -(I_1 + I_2 + I_3) \omega_1^s \\
-(I_1 - I_2 + I_3) \omega_3^s & (-I_1 + I_2 + I_3) \omega_1^s & 0
\end{bmatrix}
$$

Thus it is evident that the sum of these three matrices is skew-symmetric, although the first two matrices are not. With Eq. (88) substituted into Eq. (79), inspection of this equation and Eq. (80) permits the matrix $G'$ to be recorded in terms of 3 by 3 partitioned matrices in the expanded form shown in Fig. 6. The substitution $C \tilde{\omega}^s G = (\tilde{C}_\omega)$ has also been applied here. Note that all of the $\hat{\beta}$ terms on the right side of Eqs. (79) and (80) are accommodated in $G'$, leaving only damping terms in the symmetric matrix $D'$. The matrix $G'$ is skew-symmetric, since the transpose of the matrix in Fig. 6 is also its negative. A more compact and useful representation of this matrix can be obtained by broadening the definition of the operator tilde ($\tilde{\cdot}$). As illustrated in Eq. (44), when a tilde appears over a symbol signifying a 3 by 1 matrix, the corresponding skew-symmetric 3 by 3 matrix is implied. Henceforth, when a tilde appears over any symbol representing a column matrix that can be partitioned into 3 by 1 matrices, a skew-symmetric matrix is implied, which, as a matrix of 3 by 3 partitions, is diagonal, with the 3 by 3 skew-symmetric counterparts of the 3 by 1 elements of the column matrix ranged along the diagonal. For example, on the column matrix

$$
\Sigma_{\xi_0} \Omega^s = \begin{bmatrix} \Omega^s & 0 & \Omega^s & 0 & \cdots & \Omega^s & 0 \end{bmatrix}
$$

the tilde operator signifies

$$
(\Sigma_{\xi_0} \Omega^s)_{\tilde{\cdot}} = \begin{bmatrix}
\tilde{\omega}^s & 0 & 0 & \cdots & 0 \\
0 & 0 & 0 & \cdots & 0 \\
0 & 0 & \tilde{\Omega}^s & \cdots & 0 \\
\cdots & \cdots & \cdots & \cdots & \tilde{\Omega}^s \\
0 & 0 & 0 & \cdots & 0
\end{bmatrix}
$$

(89)
\[
G' = \begin{bmatrix}
2 [\tilde{\Omega}^1 + (\tilde{C}_n)] [m^1 - m'm'/\Omega] & 0 & -2 [\tilde{\Omega}^1 + (\tilde{C}_n)] m'm'/\Omega & 0 & \cdots & 0 \\
0 & \left(\text{tr} \, \bar{I}' \right) \bar{\Omega}^1 - 2 (\bar{I}' \bar{\omega}^1) & 0 & 0 & \cdots & 0 \\
-2 [\tilde{\Omega}^1 + (\tilde{C}_n)] m'm'/\Omega & 0 & 2 [\tilde{\Omega}^1 + (\tilde{C}_n)] [m^2 - m'm'/\Omega] & 0 & \cdots & 0 \\
0 & 0 & 0 & \left(\text{tr} \, \bar{I}' \right) \bar{\Omega}^2 - 2 (\bar{I}' \bar{\omega}^2) & \cdots & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
-2 [\tilde{\Omega}^1 + (\tilde{C}_n)] m'm'/\Omega & 0 & -2 [\tilde{\Omega}^1 + (\tilde{C}_n)] m'm'/\Omega & 0 & \cdots & 0 \\
0 & 0 & 0 & 0 & \cdots & \left(\text{tr} \, \bar{I}' \right) \bar{\Omega}^n - 2 (\bar{I}' \bar{\omega}^n) \\
\end{bmatrix}
\]

Fig. 6. Coriolis matrix
With this added convention, after inspection of Fig. 6 and Eqs. (79) and (80), the matrix $G'$ may be written in the compact form

$$G' = 2M \left( \Sigma_{E_2} \Omega^2 \right) - \Sigma_{E_2} \left[ (\vec{C}_w + \vec{C}_\theta) \Sigma_{E_2} M / \Omega \right] + M \left[ (\Sigma_{E_2} \Omega^2) + (\Sigma_{E_2} C_\theta) \right]$$

$$+ \left[ (\Sigma_{E_2} \Omega^2) + (\Sigma_{E_2} C_\theta) \right] M - M \left( \Sigma_{E_2} \Omega^2 + (\Sigma_{E_2} C_\theta) \right) \left( \Sigma_{E_2} \Omega^2 \right)$$

The matrices $A'$ and $K'$ appear in compact form in terms of the new symbol $\omega \equiv C_\theta + \Omega^2$ as

$$A' + K' = M \left( \Sigma_{E_2} \omega \right)^2 - M \Sigma_{E_2} \omega \Sigma_{E_2} M / \Omega - \left( \Sigma_{E_2} \omega \right)^2 + \left( \Sigma_{E_2} \omega \right)^2 \left( \Sigma_{E_2} \omega \right)^2 + M \left( \Sigma_{E_2} \omega \right)^2 \left( \Sigma_{E_2} \omega \right)^2 + M \left( \Sigma_{E_2} \omega \right)^2 \left( \Sigma_{E_2} \omega \right)^2 + K$$

where of particular interest is the fifth term, which is null except for the following on even rows of the main diagonal

$$- \frac{1}{2} \left( \left[ \begin{array}{c} 0 \\ (I_2 - I_2) \omega_1^2 \omega_2^2 \\
- \frac{1}{2} \omega_2 \omega_3 \\
- \frac{1}{2} \omega_2 \omega_3 
\end{array} \right] + \left[ \begin{array}{c} 0 \\ (I_2 - I_2) \omega_1^2 \omega_2^2 \\
- \frac{1}{2} \omega_2 \omega_3 \\
- \frac{1}{2} \omega_2 \omega_3 
\end{array} \right] \right)$$

and where $K$ is the stiffness matrix that defines the structural interaction forces and torques induced by deformation of the appendage. The stiffness matrix $K$ is symmetric (see Refs. 11–13), but in general there does exist a skew-symmetric matrix $A'$. Specifically, the first three terms in Eq. (91) involving angular accelerations $\omega^2$ are skew-symmetric, and the fourth such term has both symmetric and skew-symmetric parts. All matrices beyond the fifth term (expanded above) are symmetric. Figure 7 shows the first few terms in the upper left corner of the matrix $A' + K' - K$.

There remains only the explicit identification of the elements in the matrix $L'$ of Eq. (84). Compact representation is accomplished here with the new definition

$$r \equiv \left[ r^1 \ 0 \ r^2 \ 0 \ \cdots \ r^n \ 0 \right]$$

The general interpretation of the tilde operator now provides

$$\begin{bmatrix}
\tilde{r}^1 & 0 & 0 & 0 & \cdots & 0 & 0 \\
0 & 0 & 0 & 0 & \cdots & 0 & 0 \\
0 & 0 & \tilde{r}^2 & 0 & \cdots & 0 & 0 \\
0 & 0 & 0 & 0 & \cdots & 0 & 0 \\
0 & 0 & 0 & 0 & \cdots & \tilde{r}^n & 0 \\
0 & 0 & 0 & 0 & \cdots & 0 & 0
\end{bmatrix}$$

Now the 6n by 1 matrix $L'$ can be written as

$$L' = -M \Sigma_{E_2} \left( \dot{\omega} + C_\theta - \dot{\omega} + C_\theta \right) - \left( \Sigma_{E_2} \left( \Omega^2 + C_\theta \right) \right)$$

$$- M \Sigma_{E_2} C \left( \ddot{\omega} - (\tilde{R} + \tilde{c}) \dot{\omega} + \ddot{c} \Sigma_{E_2} + \ddot{\omega} \dot{\omega} \right)$$

$$+ M \left( \tilde{r} \Sigma_{E_2} \left( \dot{\omega} + C_\theta \right) - \left( \Sigma_{E_2} \Omega^2 \right) \left( \Sigma_{E_2} \Omega^2 \right) + 2 \left( \Sigma_{E_2} C_\theta \right) \left( \Sigma_{E_2} \Omega^2 \right) + \left( \Sigma_{E_2} C_\theta \right) \left( \Sigma_{E_2} \Omega^2 \right) \right)$$

$^*\text{Note from the expansion following Eq. (91) that the fifth term is symmetric if all bodies $A_1$ are spheres, or if the appendage rotates about an axis parallel to $\hat{a}_1$, $\hat{a}_2$, or $\hat{a}_3$.}$
where the new symbol $\lambda$ is a column matrix in which to store any forces or torques applied to the sub-bodies $A_1, \ldots, A_n$ other than the structural interaction forces induced by deformations. For example, field forces such as gravity forces on the appendage belong in $\lambda$, and any attitude-control jets on the appendage would contribute to $\lambda$ in a manner established by a control law.

In expanded form, the matrix $L'$ appears as a column of 3 by 1 submatrices in Fig. 8, in which the forces and torques on body $j$ in $\lambda$ are represented respectively by $f_j$ and $l_j$. This concludes the derivation of the equations of motion of an elastic appendage on a base undergoing arbitrary motion. The final result is Eq. (84), with matrices $q$, $M'$, $G'$, $K'$, $A'$, and $L'$ given in compact form by Eqs. (83), (87), (90), (91), and (94). These equations are complicated by the generality introduced with time-varying $C$, which permits application to a flexible appendage of changing attitude relative to its base (e.g., a scanning antenna). Additional complexity stems from the variable $e$, which is present to accommodate movable parts within the vehicle (other than the appendage). Specialization to the simplest configuration, with $C = E$ and $e = 0$, led to the basic Eqs. (81) and (82) for the typical submass $A_s$.

When these equations are combined as a single matrix equation such as Eq. (84), one obtains the simpler result

$$
M \left( E - \Sigma_{E_0} \Sigma_{E_0} M / \Omega \right) \ddot{q} + \{ 2M \left[ (\Sigma_{E_0} \omega)^T - \Sigma_{E_0} \overline{\omega} \Sigma_{E_0} M / \Omega \right] + M (\Sigma_{O_0} \omega)^T + (\Sigma_{O_0} \omega)^T M - [M (\Sigma_{O_0} \omega)^T + D') \dot{q} 
+ \{ (\Sigma_{O_0} \omega)^T - (M \Sigma_{O_0} \omega)^T - (M \Sigma_{O_0} \omega)^T - (M \Sigma_{O_0} \omega)^T M (\Sigma_{O_0} \omega)^T 
+ M [(\Sigma_{E_0} \omega)^T - \Sigma_{E_0} \overline{\omega} \Sigma_{E_0} M / \Omega)] + M [(\Sigma_{E_0} \omega)^T - (\Sigma_{E_0} \omega)^T - \Sigma_{E_0} \overline{\omega} \Sigma_{E_0} M / \Omega] + K \} q = 
-M \Sigma_{O_0} \dot{\omega} - (\Sigma_{O_0} \omega)^T M (\Sigma_{O_0} \omega) - M \left[ (\Sigma_{E_0} [\theta \ddot{X} - \ddot{\overline{R}} + \overline{\omega} \overline{R}] - \overline{\omega} \Sigma_{E_0} \dot{\omega} - (\Sigma_{E_0} \omega)^T (\Sigma_{E_0} \omega)^T \right] \dot{r} \}
+ \lambda
$$

Even in this case the equations remain complex in appearance. Although the specialization has greatly reduced the number of terms in the equation, no simplification of the basic structure of the equations has resulted. For the special configuration as well as the more general one, the equations of motion have the structure of Eq. (84). In both cases $M'$ is constant and symmetric; $G'$ is skew-symmetric and depending linearly on the variable and generally unknown angular velocity $\omega$ of the base $B$; $K'$ is the sum of a constant symmetric stiffness matrix $K$ and a variable symmetric matrix; and $A'$ is a variable skew-symmetric matrix comprised of terms linear in $\omega$ and terms $(T_{\alpha} - T_{\beta})(\omega_{\alpha} \omega_{\beta})$, $\alpha \neq \beta$; and $L'$ is composed of applied forces and torques, nonlinear functions of $\omega$, linear terms in $\dot{\omega}$, and the unknown mass-center motion $\ddot{X}$.

It may appear that the various matrix operators introduced here serve to obfuscate rather than clarify results, and that the appendage equations are too hopelessly large and complex for efficient simulation. Both of these impressions would be correct if the derivation were to end at this point. Many of the matrices appearing here are quite sparse, and their use in a digital computer program is rendered inefficient by the number of multiplications by zero. The number of sub-bodies $A_1, \ldots, A_n$ required for an acceptable simulation of a flexible appendage may be quite large (30 is not an unreasonably large value for $n$), and the matrix Eq. (84) is equivalent to $6n$ simultaneous coupled scalar differential equations.

Practical utility can be demonstrated for these equations only after transformation to carefully selected coordinates permits the exercise of judgment in ignoring all but a small number of variables. It is remarkable that in introducing a coordinate transformation and combining the resulting equations with appropriate equations for the balance of the vehicle, seemingly complex matrix combinations will emerge with simple physical interpretations, and coordinate truncation will permit meaningful preliminary designs of vehicle attitude-control systems to be accomplished without recourse to the computer. The results of the coordinate transformations, however, will be deferred until Section III-D so that we may first consider the remainder of the equations of motion of the vehicle.
\[ A' + K' - K = \]

\[
\begin{bmatrix}
[\tilde{u}^2 + (\tilde{C}_w) + \tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] [m^1 - m^1 m^1/\Omega^1] & 0 & -[\tilde{u}^2 + (\tilde{C}_w) + \tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] m^1 m^1/\Omega^1 \\
0 & [-\tilde{I} (\tilde{\Omega}^2) + \tilde{I} \tilde{e}^2 - \tilde{e} (\tilde{I} \tilde{e}^2) + \tilde{e} \tilde{I} \tilde{e}^2] & 0 \\
- [\tilde{u}^2 + (\tilde{C}_w) + \tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] m^1 m^1/\Omega^1 & 0 & [\tilde{u}^2 + (\tilde{C}_w) + \tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] [m^2 - m^2 m^2/\Omega^1] \\
0 & 0 & 0 \\
. & . & . \\
. & . & . \\
. & . & . \\
\end{bmatrix}
\]

where \( \epsilon^e = C_w + \Omega^1 \).

Fig. 7. Modified stiffness matrix

\[ L' = \begin{bmatrix}
-m^1 \{ C [\Theta \dot{X} - (\tilde{R} + \tilde{e}) \dot{\omega} + \dot{\omega} - 2 \tilde{C}_w \tilde{u} + \tilde{C}_w (R + e) - \tilde{r} (\tilde{u}^2 + \tilde{C}_w \tilde{u})] + [\tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] r^1 \} + f^1 \\
-\tilde{I}^1 [\tilde{u}^2 + C_w - \tilde{u}^2 C_w] - [\tilde{u}^2 + (\tilde{C}_w)] I^1 [\Omega^1 + C_w] + \tilde{I} \\
-m^2 \{ C [\Theta \dot{X} - (\tilde{R} + \tilde{e}) \dot{\omega} + \dot{\omega} - 2 \tilde{C}_w \tilde{u} + \tilde{C}_w (R + e) - \tilde{r} (\tilde{u}^2 + \tilde{C}_w \tilde{u})] + [\tilde{u}^2 \tilde{u}^2 + 2 (\tilde{C}_w) \tilde{u} + (\tilde{C}_w) (\tilde{C}_w)] r^1 \} + f^2 \\
-\tilde{I}^2 [\tilde{u}^2 + C_w - \tilde{u}^2 C_w] - [\tilde{u}^2 + (\tilde{C}_w)] I^1 [\Omega^1 + C_w] + \tilde{I} \\
. & . & . \\
. & . & . \\
. & . & . \\
-\tilde{I}^n [\tilde{u}^2 + C_w - \tilde{u}^2 C_w] - [\tilde{u}^2 + (\tilde{C}_w)] I^n [\Omega^1 + C_w] + \tilde{I}^n \\
\end{bmatrix}
\]

Fig. 8. Forcing function
C. Vehicle Equations

Immediately upon shifting attention from the appendage deformations to the time behavior of the total vehicle, the analyst confronts a basic question. Should he simply isolate the rigid body to which the appendage is attached and derive its equations of motion, treating the forces and torques applied to it by the appendage as "external"? Or should he instead write equations of motion for a dynamic system that includes both the rigid body and the flexible appendage? A parallel question was considered in Section II, in which augmented-body methods were contrasted with nested-body methods. In the context of hybrid-coordinate analysis, the answer to this question is complicated by the anticipated truncation of modal coordinates of the appendage. If the analyst isolates the body to which the appendage is attached, he must be sure that in the truncation process he retains a valid representation of the interaction forces between base and appendage. Because of this difficulty, it seems generally desirable to include the appendage with the rigid body in the dynamic system, and this will be the method used here. In certain cases, however, when the vehicle undergoes large changes in configuration, it may be most efficient computationally to treat the appendage interaction forces as external forces applied to the rigid body. Then some artifice is required (such as the "synthetic modes" of Ref. 28) to assure that coordinate truncation does not invalidate the interaction force expression. This method is explored in Section III-E of this report.

The immediate task is the derivation of equations of motion of a typical vehicle that includes flexible appendages. No attempt will be made here to establish a general set of equations applicable to virtually all space vehicles. Attention is instead restricted to a class of vehicle that may be modeled as a rigid body $B_1$ with two flexible appendages ($A_1$ and $A_2$), a balanced rigid symmetric rotor $B_3$, and a single rigid body $B_2$ attached to $B_1$ with a single degree of translational freedom.

The vehicle sketched in Fig. 3a does not quite meet the restrictions of the vehicle equations to be derived here; the rotor $B_3$ of Fig. 3a is not rigid and symmetric, since it includes an internal body $B_4$ and two flexible appendages $A_4$ and $A_5$. Figure 9 better typifies the vehicle to which the following equations apply directly. It is an idealized tri-spin satellite, designed to maintain spin stabilization with the rigid symmetric rotor $B_3$, while a control system between bodies $B_1$ and $B_3$ maintains the rigid platform $B_1$ in an earth-pointing attitude. The flexible antenna $A_1$ is attached to $B_1$ at a hinge that permits limited controlled relative motion. A solar-cell array $A_2$ is also attached to $B_1$, with an auxiliary control system maintaining the required sun-pointing orientation. A linear oscillator $B_2$ acts as a nutation damper for the satellite; its single degree of translational freedom is in a direction transverse to the bearing axis between $B_1$ and $B_3$.

Although the vehicle of Fig. 9 is of substantial interest in itself—and by specializing to omit components or fix joints one can use its equations of motion for a very wide range of space vehicles—its derivation is included here principally for its illustrative value. The analyst must be expected to approach the derivation of equations of motion of each space vehicle configuration individually, if computational efficiency is to result. The appendage equations derived in Section III-B, on the other hand, should be applied directly to any flexible appendage within the definition offered, and should not require rederivation for specific cases.

One may write in application to any material system the basic Newton-Euler equations, $F = \boldsymbol{\eta} a$ and $T = \boldsymbol{H}$, where $\boldsymbol{H}$ is the vehicle mass, $F$ is the resultant applied force, and $a$, $\boldsymbol{H}$, and $T$ are, respectively, the inertial acceleration, angular momentum, and resultant torque, all referred to the vehicle mass center.

In deriving the rotational equation from

$$T = \dot{\boldsymbol{H}}$$

(96)

Fig. 9. Tri-spin satellite (cross-sectional view)
one simply substitutes the angular momentum definition

\[ H = \int \rho \times \dot{\rho} \, dm \]  

(97)

where \( \rho \) is the generic position vector from the vehicle mass center to a typical differential mass element of the vehicle.

Define a new vector \( p \) as the generic position vector from the point \( O \) fixed in body \( B_i \) and replace \( \rho \) in Eq. (91) by \( p + \epsilon \) (recalling that \( -\epsilon \) defines the displacement in \( B_i \) of the vehicle mass center \( eM \) from its nominal location at \( O \)). The result is then differentiated (making use once more of Eq. 38) and the integral rewritten as follows, employing a convention whereby differentiation in the reference frame of \( B_i \) is signified by an overcircle (e.g., \( \dot{p} \)):

\[ H = \int (p + \epsilon) \times (\dot{p} + \dot{\epsilon}) \, dm \]

\[ = \int (p + \epsilon) \times (\dot{p} + \dot{\epsilon}) \, dm \]

\[ + \int (p + \epsilon) \times [\omega \times (p + \epsilon)] \, dm \]

\[ = \epsilon \times \int (\dot{p} + \dot{\epsilon}) \, dm + \int p \times \dot{\rho} \, dm - \epsilon \times \int p \, dm \]

\[ + \int p \, dm \times (\omega \times \epsilon) \]

\[ + \int p \times (\omega \times p) \, dm + \epsilon \times \left[ \omega \times \int (p + \epsilon) \, dm \right] \]

(98)

The mass-center definition requires

\[ \int (p + \epsilon) \, dm = 0 \]

(99)

so

\[ \int p \, dm = -\epsilon \int dm = -\epsilon \Omega \]

(100)

and

\[ \int (\dot{p} + \dot{\epsilon}) \, dm = \frac{d}{dt} \int (p + \epsilon) \, dm = 0 \]

(101)

Thus many of the integrals in Eq. (98) become either zero or a simple vector. The immediate result is

\[ H = \int p \times (\omega \times p) \, dm \]

\[ + \int p \times (\omega \times p) \, dm \]

\[ = \int p \times (\omega \times p) \, dm + \int p \times (\omega \times p) \, dm \]

(102)

The last integral in Eq. (102) may be recognized as the dot-product of the inertia dyadic \( I \) of the vehicle for point \( O \) and the inertial angular velocity of the base \( B_i \). This identification can be established in detail as follows:

\[ \int p \times (\omega \times p) \, dm = \int [p \cdot p \omega - p \cdot \omega p] \, dm \]

\[ = \int [p \cdot p \omega - p \cdot \omega p] \, dm \cdot \omega \Rightarrow I \cdot \omega \]

(103)

where \( \mathbf{E} \) is the unit dyadic (so \( \mathbf{E} \cdot \omega = \omega \)). Thus the vehicle angular momentum with respect to its mass center is

\[ H = I \cdot \omega + \Omega \hat{e} \times c + \int p \times \dot{\rho} \, dm \]

(104)

It should be noted that when \( \epsilon \) is due only to appendage deformations, assumed arbitrarily small, the product \( \Omega \hat{e} \times c \) is of second degree and therefore negligible. For the vehicle illustrated in Fig. 9, the mass center of the appendage \( A^2 \) is nominally fixed in \( B_i \), so the large relative motion permitted between these components does not contribute to \( \epsilon \). The mass of the translating body \( B_2 \) makes the contributions of this damper body to \( \epsilon \) small in an engineering sense (relatively small, but not infinitesimal), and for certain purposes (e.g., stability analysis of balanced tri-spin motion) the damper translation may be assumed arbitrarily small. Thus the contribution of \( B_2 \) to \( \Omega \hat{e} \times c \) may also be negligible. For the vehicle illustrated, only the rotation of \( A^3 \) on its hinge might contribute materially to \( \epsilon \), and even this might be only an occasional influence. The term \( \Omega \hat{e} \times c \) is preserved in this derivation for generality, but it should be recognized that it should be retained only in exceptional applications.

The rotational equation (96) requires the inertial differentiation of the angular momentum (Eq. 104), yielding
(with dyadic differentiation as in Eq. 70)

\[
T = \mathbf{1} \cdot \dot{\omega} + \mathbf{i} \cdot \dot{\omega} + \Theta \dot{\mathbf{e}} \times \mathbf{c} + \frac{\text{id}}{\text{dt}} \int \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m
\]

\[
= \mathbf{1} \cdot \dot{\omega} + \omega \times \mathbf{1} \cdot \omega + \mathbf{i} \cdot \dot{\omega} + \Theta \left[ \dot{\mathbf{e}} + 2 \omega \times \mathbf{c} + \omega \times (\omega \times \mathbf{c}) + \dot{\mathbf{c}} \right] \times \mathbf{c} + \frac{\text{id}}{\text{dt}} \int \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m
\]

(105)

Note that Euler's equations emerge here for the special case of a rigid vehicle, since then only the first two terms on the right are nonzero. Another special case of interest is that of the nominally stationary base \(B_t\), for which \(b\) is infinitesimal and Eq. (105) linearizes to

\[
T = \mathbf{1} \cdot \dot{\omega} + \int \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m
\]

(106)

The challenge of Eqs. (105) and (106) is the replacement of the integrals by more convenient functions. Relative derivatives \(\dot{\mathbf{p}}\) and \(\dot{\mathbf{p}}\) will exist only for mass elements that move relative to rigid body \(B_t\), so the separate contributions of the rotor \(B_3\), the damper \(B_2\), and the flexible appendages \(A_1\) and \(A_2\) can be calculated in turn.

For the rotor, the relevant integral is simply

\[
\frac{\text{id}}{\text{dt}} \int_{B_3} \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m = \frac{\text{id}}{\text{dt}} \int_{B_3} \mathbf{p} \times (\mathbf{\Omega} \times \mathbf{p}) \, \text{d}m = \frac{\text{id}}{\text{dt}} (\mathbf{J} \cdot \mathbf{\Omega}) = \mathbf{h}
\]

(107)

where \(\mathbf{\Omega}\) is the angular velocity of \(B_t\) relative to \(B_3\), \(\mathbf{J}\) is the inertia dyadic of the rotor, and \(\mathbf{h}\) is the angular momentum of the rotor \(B_3\) relative to the base \(B_t\).

For the translating body \(B_2\), the integration is trivial, since all parts of \(B_2\) have the same velocity \(\dot{\mathbf{b}}\) in \(B_1\). Thus if \(m\) is the mass of \(B_2\), and its position relative to \(O\) is given in the \(B_1\)-fixed vector basis \(\mathbf{b}_1, \mathbf{b}_2, \mathbf{b}_3\), by \(\mathbf{b}_b + \xi\mathbf{b}_1,

\[
\int_{A} \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m = \int_{A} (R + r^* + u^* + \rho^*) \times [\mathbf{\Omega}^* \times (r^* + u^*) + (a)^* \dot{u}^* + (a)^* \dot{\beta}^*] \times \rho^* \, \text{d}m
\]

\[
= R \times (a)^* \int_{A} \dot{u}^* \, \text{d}m + \int_{A} r^* \times (a)^* \dot{u}^* \, \text{d}m + R \times \left[ \mathbf{\Omega}^* \times \int_{A} (r^* + u^*) \, \text{d}m \right]
\]

\[
+ \int_{A} r^* \times [\mathbf{\Omega}^* \times (r^* + u^*)] \, \text{d}m + \int_{A} u^* \times (\mathbf{\Omega}^* \times r^*) \, \text{d}m + \int_{A} \rho^* \times [(\mathbf{\Omega}^* + (a)^* \dot{\beta}^*) \times \rho^*] \, \text{d}m
\]

(108)

since in every term the integral over \(A\) can be replaced by the sum of \(n\) integrals over the individual sub-bodies \(A_1, \ldots, A_n\), and

\[
\int_{A} \rho^* \, \text{d}m = 0
\]

Finally, the contributions of the appendages to this integral must be sought. Return for this purpose to the concept of the discrete parameter model of the appendage, and the definitions of \(R, r^*, \) and \(u^*\) in Eq. (36) and Fig. 4. In addition, let \(\rho^s\) represent a generic position vector from the mass center \(P_s\) of sub-body \(A_s\) to a typical differential mass element in \(A_s\). The appropriate integral then becomes, for a typical appendage \(A\) (here either \(A^1\) or \(A^2\))

\[
\int_{A} \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m = \int_{A} (R + r^* + u^* + \rho^s) \times (\dot{r}^s + \dot{u}^s + \dot{\beta}^s) \, \text{d}m
\]

(109)

Yet another application of the vector differentiation formula in Eq. (38) is required for the interpretation of Eq. (109), and this can be done conveniently by writing the individual deformation vectors in terms of appropriate vector arrays (as in Eqs. 37 and 72). Recalling that \(\mathbf{\Omega}^s\) is the angular velocity of \(\{a\}\) relative to \(\{b\}\) for a given appendage \(A\), and \((a)^* \dot{\beta}^s\) is the angular velocity of \(A_s\) relative to \(A\), one can write Eq. (109) as

\[
\int_{A} \mathbf{p} \times \dot{\mathbf{p}} \, \text{d}m = \int_{A} (R + r^* + u^* + \rho^s) \times (\dot{r}^s + \dot{u}^s + \dot{\beta}^s) \, \text{d}m
\]

(110)
This result, when substituted into Eq. (105), would be appropriate for the simulation of the vehicle of Fig. 9 in the unlikely mode of motion in which the appendages are undergoing gradual change of orientation with respect to (however artificially) for equations such as Eq. (105), linearized in deformations only. With linearization in only, and deformation velocities and accelerations probably exceed, in most space vehicles, those due to prescribed o

\[ \int \sigma \, dm = m_s \]

Thus that equation becomes (with \( \Sigma \) denoting the sum over \( s \) ranging from 1 to \( n \))

\[ \int A \, p \times \dot{p} \, dm = R \times (a)^T \Sigma m_s \, \dot{u}^s + \Sigma r^s \times (a)^T m_s \, \dot{u}^s \]

\[ + R \times [\Omega^s \times \Sigma m_s \,(r^s + u^s)] + \Sigma m_s \, r^s \times (\Omega^s \times (r^s + u^s)) + \Sigma m_s \, u^s \times (\Omega^s \times r^s) + \Sigma \, \dot{r} \cdot ((a)^T \dot{\beta}^s + \Omega^s) \]

Equation (105) requires the inertial derivative of Eq. (111), which is

\[ \frac{d}{dt} \int A \, p \times \dot{p} \, dm = (\omega \times R) \times (a)^T \Sigma m_s \, \dot{u}^s + R \times [(a)^T \Sigma m_s \, \dot{u}^s + (\omega + \Omega^s) \times (a)^T \Sigma m_s \, \dot{u}^s] \]

\[ + \Sigma [(\omega + \Omega^s) \times r^s] \times (a)^T m_s \, \dot{u}^s + \Sigma r^s \times [(a)^T m_s \, \dot{u}^s + (\omega + \Omega^s) \times (a)^T m_s \, \dot{u}^s] \]

\[ + (\omega \times R) \times [\Omega^s \times \Sigma m_s \,(r^s + u^s)] + R \times [\dot{\Omega}^s + \omega \times \Omega^s] \times \Sigma m_s \,(r^s + u^s)] \]

\[ + R \times [\Omega^s \times \Sigma m_s \,(a)^T \dot{u}^s] + R \times (\Omega^s \times \Sigma m_s [(\omega + \Omega^s) \times (r^s + u^s)]) \]

\[ + \Sigma m_s \,(\omega \times \Omega^s \times r^s) \times [\Omega^s \times (r^s + u^s)] + \Sigma m_s \, r^s \times [(\dot{\Omega}^s + \omega \times \Omega^s \times (r^s + u^s]) \]

\[ + \Sigma m_s \, u^s \times [(\dot{\Omega}^s + \omega \times \Omega^s \times r^s)] + \Sigma m_s \, u^s \times (\Omega^s \times [\omega \times \Omega^s \times r^s]) \]

\[ + \Sigma \, \dot{r} \cdot (a)^T \dot{\beta}^s + (\omega + \Omega^s) \times (a)^T \dot{\beta}^s] + (\omega + \Omega^s) \times \dot{r} \cdot (a)^T \dot{\beta}^s - \dot{r} \times \omega \times (\omega + \Omega^s) \cdot (a)^T \dot{\beta}^s \]

\[ + \Sigma \, \dot{r} \times (\dot{\Omega}^s + \omega \times \Omega^s) + \Sigma [(\omega + \Omega^s \times (a)^T \dot{\beta}^s)] \times \dot{r} \times \Omega^s - \Sigma \, \dot{r} \times (\omega + (a)^T \dot{\beta}^s) \times \Omega^s \]

Equation (112) becomes

\[ \frac{d}{dt} \int A \, p \times \dot{p} \, dm = (\omega \times R) \times (a)^T \Sigma m_s \, \dot{u}^s + R \times [(a)^T \Sigma m_s \, \dot{u}^s + \omega \times (a)^T \Sigma m_s \, \dot{u}^s] \]

\[ + \Sigma (\omega \times r^s) \times (a)^T m_s \, \dot{u}^s + \Sigma r^s \times [(a)^T m_s \, \dot{u}^s + \omega \times (a)^T m_s \, \dot{u}^s] \]

\[ + (\omega \times R) \times (\Omega^s \times \Sigma m_s \, r^s) + R \times (\dot{\Omega}^s + \omega \times \Omega^s) \times \Sigma m_s \, r^s + R \times [\Omega^s \times \Sigma m_s \,(\omega \times r^s)] \]

\[ + \Sigma m_s \, r^s \times [(\dot{\Omega}^s + \omega \times \Omega^s \times r^s)] + \Sigma m_s \, r^s \times (\Omega^s \times (\omega \times r^s)] + \Sigma \, \dot{r} \times (\dot{\Omega}^s + \omega \times \Omega^s) \]

\[ + \Sigma [\omega \times \dot{r} \times \Omega^s - \dot{r} \times \omega \times \Omega^s] + \Sigma \, \dot{r} \times [(a)^T \dot{\beta}^s + \omega \times (a)^T \dot{\beta}^s] + \omega \times \dot{r} \times (a)^T \dot{\beta}^s - \dot{r} \times \omega \times (a)^T \dot{\beta}^s \]

This unwieldy expression, repeated for each appendage attached to body \( B_i \), must together with Eqs. (107) and (108) be substituted into the rotational equation of motion (Eq. (105)) in the general case of arbitrary base motion. Furthermore, the contribution of appendage deformation to \( i \) in Eq. (105) would have to be calculated. In most engineering applications, this degree of complexity is unwarranted. In the foregoing, only appendage deformations have been restricted, and these have been assumed to remain infinitesimal. In fact, they are at best small in an engineering sense only, and deformation velocities and accelerations probably exceed, in most space vehicles, those due to prescribed relative motions (such as \( \Omega^s \) and \( \dot{\Omega}^s \)). It is thus most reasonable in an engineering application to linearize in these latter variables also, although it should be understood that the resulting equations lack the rigor that can be claimed (however artificially) for equations such as Eq. (105), linearized in deformations only. With linearization in \( \Omega^s \) and \( \dot{\Omega}^s \), Eq. (112) becomes

\[ \frac{d}{dt} \int A \, p \times \dot{p} \, dm = (\omega \times R) \times (a)^T \Sigma m_s \, \dot{u}^s + R \times [(a)^T \Sigma m_s \, \dot{u}^s + \omega \times (a)^T \Sigma m_s \, \dot{u}^s] \]

\[ + \Sigma (\omega \times r^s) \times (a)^T m_s \, \dot{u}^s + \Sigma r^s \times [(a)^T m_s \, \dot{u}^s + \omega \times (a)^T m_s \, \dot{u}^s] \]

\[ + (\omega \times R) \times (\Omega^s \times \Sigma m_s \, r^s) + R \times (\dot{\Omega}^s + \omega \times \Omega^s) \times \Sigma m_s \, r^s + R \times [\Omega^s \times \Sigma m_s \,(\omega \times r^s)] \]

\[ + \Sigma m_s \, r^s \times [(\dot{\Omega}^s + \omega \times \Omega^s \times r^s)] + \Sigma m_s \, r^s \times (\Omega^s \times (\omega \times r^s)] + \Sigma \, \dot{r} \times (\dot{\Omega}^s + \omega \times \Omega^s) \]

\[ + \Sigma [\omega \times \dot{r} \times \Omega^s - \dot{r} \times \omega \times \Omega^s] + \Sigma \, \dot{r} \times [(a)^T \dot{\beta}^s + \omega \times (a)^T \dot{\beta}^s] + \omega \times \dot{r} \times (a)^T \dot{\beta}^s - \dot{r} \times \omega \times (a)^T \dot{\beta}^s \]

This result, when substituted into Eq. (105), would be appropriate for the simulation of the vehicle of Fig. 9 in the unlikely mode of motion in which the appendages are undergoing gradual change of orientation with respect to \( B_i \),
while the inertial angular velocity $\omega$ of $B_1$ is large. More useful forms of this result appear either when $\Omega^e$ is identically zero, which leaves

$$\frac{d}{dt} \int p \times \dot{p} \, dm = (\omega \times R) \times (a) \Sigma m_s \hat{u}^t + R \times [(a) \Sigma m_s \hat{u}^t + \omega \times (a) \Sigma m_s \hat{u}^t]$$

$$+ \Sigma (\omega \times r^t) \times (a) \Sigma m_s \hat{u}^t + \Sigma r^t \times [(a) \Sigma m_s \hat{u}^t + \omega \times (a) \Sigma m_s \hat{u}^t]$$

$$+ \Sigma I^* \cdot [(a) \Sigma \hat{\beta}^t + \omega \times (a) \Sigma \hat{\beta}^t] + \omega \times \Sigma I^* \cdot (a) \Sigma \hat{\beta}^t - \Sigma I^* \times \omega \cdot (a) \Sigma \hat{\beta}^t$$

(114)

or when $\omega$ is also assumed to be small and included in the linearization, with the result

$$\frac{d}{dt} \int p \times \dot{p} \, dm = R \times (a) \Sigma m_s \hat{u}^t + \Sigma r^t \times (a) \Sigma m_s \hat{u}^t + R \times (\hat{\Omega}^e \times \Sigma m_s, r^t) + \Sigma m_s, r^t \times (\hat{\Omega}^e \times r^t) + \Sigma I^* \cdot (a) \Sigma \hat{\beta}^t$$

$$+ \Sigma I^* \cdot \hat{\Omega}^e$$

(115)

These two special cases may be of value in practical applications, since they accommodate the tri-spin vehicle with flexible appendages on the de-spun body, and also the dual-spin vehicle with appendages on the spinning body. Even further specialization may be warranted, since spinning vehicles with flexible appendages are usually simple "spinners," rather than dual-spin vehicles, and a de-spun body with flexible appendages is more apt to be part of a dual-spin vehicle than of a tri-spin vehicle, in which case the relative motion $\Omega^e$ is usually zero.

The essential differences among the expanded forms of the rotational equation (105) are simply differences in the size of the equations and the amount of bookkeeping involved in writing them in matrix form. Because the purpose here is simply to illustrate the structure of the vehicle equations, attention is restricted henceforth to a dual-spin vehicle with a rigid, symmetric rotor, a nutation damper, and a single flexible appendage without rotational capability relative to its base ($\Omega^e = 0$ and $C = E$ so $(a) = (b)$). The platform angular velocity $\omega$ is unrestricted, so the equations to follow may easily be specialized either to the simple spinner with flexible appendages (by omitting the rotor) or to the dual-spin vehicle with an appendage on a de-spun body (by linearizing in $\omega$). By taking both of these steps (omitting the rotor and linearizing in $\omega$), one obtains equations of motion of a space vehicle with three-axis active reaction-jet control in an inertially stationary nominal orientation.

By inspection of Eqs. (105), (107), (108), and (114), the required vector-dyadic equation is obtained as follows:

$$T = I \cdot \omega + \omega \times I \cdot \omega + \dot{I} \cdot \omega + \dot{I} \cdot \dot{b} + mb (\xi b_2 + \xi \omega \times b_2) + (\omega \times R) \times (b)^T \Sigma m_s \hat{u}^t$$

$$+ R \times [(b)^T \Sigma m_s \hat{u}^t + \omega \times (b)^T \Sigma m_s \hat{u}^t] + \Sigma (\omega \times r^t) \times (b)^T m_s \hat{u}^t + \Sigma r^t \times [(b)^T m_s \hat{u}^t + \omega \times (b)^T m_s \hat{u}^t]$$

$$+ \Sigma I^* \cdot [(b)^T \hat{\beta}^t + \omega \times (b)^T \hat{\beta}^t] + \omega \times \Sigma I^* \cdot (b)^T \hat{\beta}^t - \Sigma I^* \times \omega \cdot (b)^T \hat{\beta}^t$$

(116)

Note that the mass-center motion $c$ has been assumed to be infinitesimal, in accordance with the indicated specialization.

Terms involving $I$ in Eq. (116) are time-variable, since the inertia dyadic of the composite vehicle depends on its deformations. By definition (see Eq. 103), the vehicle inertia dyadic for point $O$ is given by

$$I = \int (p \cdot p E - pp) \, dm$$

(117)

Let the generic symbol $p^*$ be the value of $p$ when the vehicle is undeformed, so that the inertia dyadic of the undeformed vehicle is

$$I^* = \int (p^* \cdot p^* E - p^* p^*) \, dm$$

(118)

For all points of the vehicle except those in the flexible appendage and the nutation damper mass, $p$ is identically $p^*$. For points in the translating damper body $B_2$,

$$p = p^* + \xi b_1 = bb_3 + \xi b_1$$

(119)
and for a typical point of body $A_v$ of the appendage,

$$p = p^s + u^s + p^s = R + r^s + u^s + p^s \quad (120)$$

In addition to the contribution of the translational deformation $u^s$ to the deformed vehicle inertia dyadic $I$, there is a contribution from the rotational deformation $\beta^s$ for each sub-body $A_v$. If the vector array $\{a^s\}$ consists of a dextral orthonormal set fixed in $A_v$, then the inertia dyadic $I^r$ of $A_v$ referenced to the mass center $P_v$ may be written in the $\{a^s\}$ basis as

$$I^r = \{a^s\}^T I^s \{a^s\} \quad (121)$$

If now $\{a^s\}$ is aligned with $\{b\}$ when the appendage is undeformed, then the deformation $\{b\}^T \beta^s$ yields the unit vector relationship

$$\{a^s\} = (E - \beta^s) \{b\} \quad (122)$$

since $E - \beta^s$ is the linear approximation of the direction cosine matrix generated by the orthogonal 1-2-3-type rotations $\beta_1, \beta_2, \beta_3$. Thus in terms of the vector basis $\{b\}$, the inertia dyadic for body $A_v$ becomes

$$I^r = \{b\}^T (E - \beta^s) I^s (E - \beta^s) \{b\}^s$$

$$= \{b\}^T (E + \beta^s) I^s (E - \beta^s) \{b\}^r$$

$$= \{b\}^T \{b\} + \{b\}^T (\beta^s I^s - I^s \beta^s) \{b\} \quad (123)$$

The second term is therefore the contribution of rotation $\beta^s$ to the inertia dyadic $I$ of the vehicle.

From Eqs. (117) through (120) and (123), it follows that the vehicle inertia dyadic is

$$I = I^r - mb\zeta (b, b_1 + b, b_3) + \Sigma m_3 [2 (R + r^s) \cdot u^s E - (R + r^s) u^s - u^s (R + r^s)] + \Sigma (b)^r (\beta^s I^s - I^s \beta^s) \{b\} \quad (124)$$

Noting that the identity dyadic is

$$E = \{b\}^s \{b\} \quad (125)$$

so that, for example,

$$R \cdot u^s E = R^T \{b\} \cdot \{b\}^s u^s \{b\} \{b\} = R^T u^s \{b\} \{b\} = \{b\}^r R^T u^s \{b\}$$

one may write the inertia dyadic as (defining $E^a$ as the $a^{th}$ column of the 3 by 3 matrix $E$)

$$I = I^r - \{b\}^s mb\zeta (E^a E^a + E^a E^a) \{b\}$$

$$+ \{b\}^s \Sigma m_3 [2 (R + r^s) \cdot u^s E - (R + r^s) u^s - u^s (R + r^s)] \{b\} + \{b\}^r \Sigma (\beta^s I^s - I^s \beta^s) \{b\} \quad (126)$$

The derivative $\dot{I}$ is readily obtained from Eq. (126) as

$$\dot{I} = \{b\}^s \{ -mb\zeta (E^a E^a + E^a E^a) + \Sigma m_3 [2 (R + r^s) \dot{u}^s E - (R + r^s) \dot{u}^s - \dot{u}^s (R + r^s)] + \Sigma (\beta^s I^s - I^s \beta^s) \} \{b\} \quad (127)$$

Finally, the complete and explicit vector-dyadic equation of rotational motion can be obtained by substituting Eqs. (126) and (127) into Eq. (116). In the process, all vectors and dyadics are written in terms of the vector array $\{b\}$, and the matrix representation of cross-products is employed (see Eq. 44). The result is

$$\{b\}^r T = \{b\}^s \{I^* \omega - mb\zeta (E^a E^a + E^a E^a) \omega + \Sigma m_3 [2 (R + r^s) \omega E - (R + r^s) \omega - \omega (R + r^s)] \omega + \Sigma (\beta^s I^s - I^s \beta^s) \omega$$

$$+ \Sigma (\beta^s I^s - I^s \beta^s) \omega - mb\zeta (E^a E^a + E^a E^a) \omega + \Sigma m_3 [2 (R + r^s) \dot{u}^s E - (R + r^s) \dot{u}^s - \dot{u}^s (R + r^s)] \omega$$

$$+ \Sigma (\beta^s I^s - I^s \beta^s) \omega + \dot{h} + m \dot{b}^2 \dot{E}^2 + mb\zeta \dot{E}^2 + \Sigma \dot{m} \dot{E} \dot{u}^s + \Sigma \dot{m} \dot{u} \dot{u} + \Sigma \dot{m} \dot{u} \dot{u} + \Sigma \dot{m} \dot{u} \dot{u} + \Sigma \dot{m} \dot{u} \dot{u}$$

$$+ \Sigma \dot{m} \dot{u} \dot{u} + \Sigma \dot{m} \dot{u} \dot{u} + \Sigma \dot{m} \dot{u} \dot{u} - \Sigma \dot{m} \dot{u} \dot{u} \quad (128)$$
Symbols \( T, h \), and \( I^* \) are new, being defined here by their context. The identities

\[
P \times \omega \cdot (b)T \dot{\beta} = P \cdot \omega \times (b)T \dot{\beta} = (b)T \dot{\gamma} \wedge \dot{b}T
\]

and

\[
\dot{h} = \dot{h} + \omega \times h = (b)T \dot{h} + (b)T \dot{\omega} h
\]

have been employed in writing Eq. (128).

The matrix equation isomorphic to the vector equation (128) can be written, by inspection, simply by dropping the vector arrays (or by pre-dot-multiplying by \( b^T \)). This matrix equation is more useful when written in terms of the total \( 6n \) by \( 1 \) deformation matrix \( q \) (defined in Eq. 83), making use of the operators \( \Sigma_{OE} \) and \( \Sigma_{OF} \) (see Eq. 85). The result then adopts the form

\[
T = I^* \dot{\theta} + \dot{h} - \dot{h} \dot{\omega} + \Sigma_{OE} \dot{M}q + \dot{R} \Sigma_{EO} \dot{M}q + \Sigma_{EO} \dot{M}q + \Sigma_{OE} \dot{M}q + \Sigma_{OE} \dot{M}q
\]

where the operator \( \dot{\Sigma} \) reassembles the three-by-one submatrices of a column matrix into a three-row matrix as illustrated by \( r = [r^T \ 0 \ 0 \cdots r^T \ 0] \).

The more general Eq. (129) has the major added term \( \dot{\Sigma} \omega \), which introduces the "Euler coupling" or "gyroscopic coupling" of the vehicle as a spinning rigid body. In addition, Eq. (129) includes a multitude of nonlinear terms that reflect the influence of the vehicle deformability. They can be categorized as various kinds of acceleration terms (centripetal, coriolis, etc.), but perhaps they are best understood by reviewing their origins in the vector-dyadic equation (116).

The derivation of this equation has been the principal objective in this section. Interpretation may be facilitated by restricting attention to the special case of a dual-spin vehicle with an appendage on a nominally de-spun platform. This case is significant from an applications viewpoint, and yet the assumption that \( \omega \) is infinitesimal renders Eq. (129) quite interpretable. Since \( \omega \) is small, it can be replaced by the matrix \( \delta \), where \( \delta = [\delta_1 \ \delta_2 \ \delta_3]^T \).

The result is simply the linear equation

\[
T = I^* \delta + \dot{h} - \dot{h} \dot{\omega} + \Sigma_{EO} \dot{M}q + \dot{R} \Sigma_{EO} \dot{M}q + \Sigma_{EO} \dot{M}q + \Sigma_{EO} \dot{M}q
\]

Since \( I^* \) is the inertia matrix of the total vehicle as a rigid body, the equation \( T = I^* \delta \) would suffice if the rotor were fixed (nonrotating) and the appendage were rigid. The torque applied to the body \( B_1 \) in accelerating the relative angular rate of the rotor is \( -\dot{h} \delta \), and the "gyroscopic stiffness" supplied by the rotor is in \(-\dot{h} \delta \). The term \( mb^2 \dot{E}^2 \) reflects the "inertial torque" due to damper mass acceleration, and the last three terms reflect the effects of appendage deformation.
If $F$ is written in terms of the body-fixed vector basis $\{b\}$, related to $\{i\}$ as in Eq. (53), this equation becomes

$$(b)^T F = \mathcal{M} (i)^T \ddot{X} = \mathcal{M} (b)^T \ddot{X}$$

or in matrix terms

$$F = \mathcal{M} \ddot{X}$$  \hspace{2cm} (131)

Newton's second law also provides the equation of motion of the damped linear oscillation representing the nutation damper, except that only the $b_l$ component of the vector equation of translation is required. The vector result is

$$M (\ddot{X} + \dddot{b}_a + \dddot{b}_i + \dddot{b}_r) \cdot b_i = F^{n_l} \cdot b_i = -k \dddot{e} - d \dddot{e}$$  \hspace{2cm} (132)

where $k$ and $d$ are scalar constants of the spring and dashpot, respectively. The mass-center displacement vector $c$ is available from Eq. (60) as

$$c = -(b)^T \Sigma_{E^0} \mu q - (m \dddot{e} / \mathcal{M}) b_i = -(b)^T [\Sigma_{E^0} \mu q + m \dddot{e} / \mathcal{M}]$$

so that

$$\dddot{c} = -(b)^T [\Sigma_{E^0} \mu q + m \dddot{e} / \mathcal{M}]$$

Similar operations on $b_a = (b)^T E^3$ and $b_i = (b)^T E^1$ permit the expansion of the term in parentheses in Eq. (132), and the $b_l$ dot-product then provides the scalar equation

$$m (1 - m / \mathcal{M}) \dddot{e} + [d + 2mE^7 \Sigma_{E^1} (1 - m / \mathcal{M})] \dddot{e} + [k \dddot{e} / \Sigma_{E^0} \mu q + m \dddot{e} / \mathcal{M}]$$

where $\tau$ is the magnitude of the torque applied to the rotor along its axis of symmetry. In general $\tau$ is some combination of bearing friction and the motor torque established by a control law. Complete simulation would of course require an auxiliary equation for that control law, and except for very preliminary calculations the control law would be nonlinear.

In summary, Eqs. (136), (134), (131), (129), and (95) constitute a complete set of dynamic equations for the system. The unknowns are included in $\phi, \xi, X, \Theta, \phi, \psi, \varphi, \omega, \phi, \tau, \phi, \theta, \phi, \rho$, and $\phi$. (Equation 135 must be substituted into Eq. 129 before collecting these equations, in order to eliminate $h$.) To complete the simulation, a set of kinematic equations relating $\phi$ and $\psi$ must be included. The particular equations used will depend on the attitude parameters chosen to define the orientation of $B_i$ in inertial space (e.g., direction cosines, Euler parameters, attitude angles, etc.). These equations are available from the fundamental kinematic relationship (Ref. 16)

$$\ddot{\omega} = \dot{\omega} \dot{\phi} - \dot{\phi} \dot{\omega}$$  \hspace{2cm} (137)

but more suitable forms of the kinematic equations can be obtained for specific parameter choices (e.g., see Ref. 23).
It is then conceptually possible to collect the five dy-
namic equations (136), (134), (131), (129), and (95) to-
gether with kinematic equations from Eq. (137) and
control laws for the rotor torque \( \tau \) and the vehicle torque
\( T \), and with this set, to accomplish the simulation of a
dual-spin vehicle with a flexible appendage and a nuta-
tion damper attached to a base undergoing arbitrary
motion. By specializing these equations, one could simu-
late a simple spin-stabilized vehicle, or a vehicle with
three-axis reaction-jet control. By generalizing these equa-
tions, proceeding from vector-dyadic equations developed
here in detail, one could with straightforward labor simi-
larly simulate a vehicle controlled by several rotors
(“momentum wheels”) and having articulated elastic ap-
pendages capable of substantial relative motion. All of
this is, however, only conceptually possible. The complete
set of equations is available, but if they are to be applied
to a real space vehicle represented by a reasonably accu-
rate mathematical model, the dimension of the resulting
equations is prohibitively high, even for machine com-
putation. Remember that Eq. (95) alone consists, for
practical problems, of hundreds of second-order scalar
differential equations, and that the entire system of equa-
tions is nonlinear and highly coupled. If these equations
are to have practical value, they must be subjected to
coordinate transformations that accomplish substantial
uncoupling of the equations and permit the analyst to
exercise judgment in restricting his attention to a subset
of the present set of perhaps hundreds of coordinates, in
order to justify working with a more reasonable number
of differential equations in the simulation of the vehicle.

D. Coordinate Transformations

1. Transformation rationale. The objective of this sec-
tion is to investigate the possibility of simplifying the
equations of motion by introducing linear transformations
for some of the variables. The transformations are applied
to the appendage deformation matrix \( q \) only, leaving the
discrete coordinates of the base \( B_i \), the damper \( B_j \), and
the rotor \( B_k \) unchanged. The more remote possibility of
finding a transformation that may fruitfully be applied to
the entire system of coordinates is treated in Section IV.

Of course, an infinite variety of coordinate transforma-
tions may be devised, even within the linear class (to
which attention is presently restricted). Any transforma-
tion that provides a one-to-one mapping from one coor-
dinate system to the next yields equations that could
theoretically provide a correct simulation, but obviously
not all such transformations are advantageous. It may be,
however, that more than one useful transformation will
be available, and then the appropriate choice may depend
on individual standards of utility.

The basic difficulty in using the equations of motion
derived earlier (Eqs. 95, 129, 131, 134, and 136 or their
equivalent) is simply their dimension. A certain amount
of nonlinearity and coupling of equations may be un-
avoidable. Therefore the primary test of the utility of a
given coordinate transformation is the degree to which
it permits the truncation of the coordinate matrix, and
the consequent reduction in the dimension of the system
of equations.

Coordinate truncation is never a completely rigorous
process, since it results in an incomplete and imprecise
indication of the response of the mathematical model to
dynamic environment. As a practical matter, however,
it must be recognized that the mathematical model is but
an imperfect approximation of the vehicle being studied,
so it is sophistry to argue that coordinate truncation nec-
essarily degrades the simulation of a real vehicle. Some
degree of truncation may be appropriate even when the
most realistic efficient simulation is sought.

There are no entirely satisfactory analytical procedures
for determining the degree to which a given matrix of
coordinates can safely be truncated, nor can the relative
acceptability of truncation of alternative coordinate sys-
tems be unequivocally established. It should be noted,
however, that when a coordinate transformation pro-
vides a completely uncoupled system of scalar differential
equations, one can obtain the solution of these equations
precisely for each scalar coordinate in turn, without con-
sidering the influence of other coordinates. Therefore one
can truncate the coordinate matrix without sacrificing
the rigorous validity of the solutions for the coordinates
retained. The determination of the number of such coor-
dinates to retain for acceptable simulation is still a matter
of engineering judgment, but the validity of the solution
for those retained is related directly to the degree to
which the transformed equations are uncoupled. Accord-
ingly, the desirability of a given coordinate transforma-
tion is measured here in terms of the degree to which it
uncouples the system of differential equations employed
in the vehicle simulation.

The total system of equations required for the vehicle
simulation includes some equations in which the append-
age deformation terms are of central importance (e.g., the
appendage equation 95), some equations in which terms
in \( q \) are generally of secondary importance (e.g., the nuta-
tion damper equation 134), in which \( q \) is always multi-

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plied by the small constant matrix $\rho$, and some equations in which $q$ does not appear at all (e.g., the rotor equation 136). In the quest for a transformation that to some degree uncouples the equations, it is therefore reasonable to concentrate on a subset of the system of differential equations, excluding from consideration the equations of motion of rotor and damper, the kinematic equations relating $\phi$ and $\phi$, and the control system equations. (Note that the last of these will include $q$ when sensors or thrusters are mounted on flexible appendages.) This decision to concentrate on certain equations clearly marks a compromise with the objective of securing uncoupled equations, and, in fact, this objective became unrealizable with the announced restriction in this section to transformations of the appendage deformation coordinates $q$ only. The final equations must certainly remain coupled by the discrete coordinates of $B_1$, $B_2$, and $B_3$, and by ignoring certain equations while transformations are sought, one accepts as well the likelihood that these equations will, after transformation, remain coupled in the new deformation variables.

Attention is now restricted to Eqs. (95), (129), and (131), which are respectively the appendage equation, the vehicle rotation equation, and the vehicle translation equation. The last of these is trivial, permitting the solution

$$\theta \ddot{X} = F/\Omega \lambda$$

(138)

This substitution into Eq. (95) with the substitution $\ddot{\phi} = -m\dddot{E}^T/\Omega \lambda$ to accommodate the damper, provides the revised appendage equation

$$M (E - \Sigma_{E0} \Sigma_{E0}^T M/\Omega \lambda) \ddot{q} + 2 (M (\Sigma_{E0} \omega)^{-1} - \Sigma_{E0} \Sigma_{E0}^T M/\Omega \lambda) + M (\Sigma_{E0} \omega)^{-1} + (\Sigma_{E0} \omega)^{-1} M (\Sigma_{E0} \omega)^{-1} + D') \ddot{q}$$

$$+ (M (\Sigma_{E0} \omega)^{-1} - (\Sigma_{E0} \omega)^{-1} (\Sigma_{E0} \omega)^{-1} + (\Sigma_{E0} \omega)^{-1} M (\Sigma_{E0} \omega)^{-1}) + M [(\Sigma_{E0} \omega)^{-1} - \Sigma_{E0} \Sigma_{E0}^T M/\Omega \lambda] + M [(\Sigma_{E0} \omega)^{-1} - (\Sigma_{E0} \omega)^{-1} (\Sigma_{E0} \omega)^{-1} M (\Sigma_{E0} \omega)^{-1})$$

$$- M (\Sigma_{E0} - \Sigma_{E0} \tilde{R} - \gamma \Sigma_{E0}) \phi - M (\Sigma_{E0} \Sigma_{E0} R - (\Sigma_{E0} \omega)^{-1} (\Sigma_{E0} \omega)^{-1} \tilde{R}) - (\Sigma_{E0} \omega)^{-1} M (\Sigma_{E0} \omega)^{-1}$$

$$- M \Sigma_{E0} F/\Omega \lambda + M \Sigma_{E0} m \dddot{E}^T/\Omega \lambda + \lambda$$

(139)

This equation consists of 6n second-order scalar equations and can be written as a matrix equation of the structure of Eq. (84), or somewhat more explicitly as

$$M' \dddot{q} + D' \dddot{q} + C' \dddot{q} + A' \dddot{q} = L' = -M (\Sigma_{E0} - \Sigma_{E0} \tilde{R} - \gamma \Sigma_{E0}) \phi + N^c - M \Sigma_{E0} F/\Omega \lambda + M \Sigma_{E0} m \dddot{E}^T/\Omega \lambda + \lambda$$

(140)

where $N^c$ represents the nonlinear terms in $\omega$ due to centrifugal forces on the appendage. Here $G'$ depends linearly on $\omega$, $A'$ and $K'$ depend linearly on $\phi$ and non-linearly on $\omega$ (involving only second-degree terms). As noted in the discussion of Eq. (84), $M'$, $D'$, and $K'$ are symmetric and $G'$ and $A'$ are skew-symmetric.

The remaining equation of primary interest, Eq. (129), consists of only three scalar equations, although many of its terms involve matrix operators that accomplish summations over 6n elements. The appearance of the variable $q$ and its derivatives in Eq. (129) is again linear, and again $\phi$ appears only in the first power and $\omega$ in the second, so it would be possible to write this equation in the form of Eq. (84). This would be misleading, however, since the coefficient matrices would be rectangular (3 by 6n), and the role of Eq. (129) as the vehicle rotational equation would be obscured. To emphasize the primary physical significance of this equation, it is rewritten in symbolic form as

$$I \dddot{\phi} + 2 \Sigma_{E0} (I + G - \tilde{R}) \phi =$$

$$- (\Sigma_{E0}^T + \tilde{R} \Sigma_{E0}^T + \Sigma_{E0} \tilde{R}) M \dddot{q} - \tilde{h} - m \dddot{E}^T + T$$

(141)

This form of the vehicle rotational equation can be constructed from Eq. (129) quite mechanically by defining $I$ as the coefficient matrix of $\phi$, observing that this same matrix permits $\Sigma_{E0}$ to accommodate all second-degree terms in $\omega$, and then defining $G$ to include all terms in the coefficient matrix of $\omega$ not absorbed by $\tilde{I}$ or $\tilde{R}$.

Much more insight into the significance of $I$ and $G$ may be gained by inspecting Eq. (116), which is the vector-zydric antecedent of Eq. (129). With this examination comes the realization that $I$ is the time-variable inertia matrix of the total vehicle in basis $\{b\}$, referred to the vehicle mass center. The term $G_{\omega}$ accommodates the "inertial torques" applied to the vehicle due to the coriolis accelerations of appendage and damper, so it has the same physical origins as the matrix $G'$ of Eq. (140). It should be noted that
the coefficient of \( \dot{q} \) in Eq. (141) is the transpose of the coefficient of \( \dot{\omega} \) in Eq. (140), i.e.,

\[
[(\Sigma_{\omega e}^{T} + \bar{R} \Sigma_{\omega e} + \Sigma_{\omega e}^{T}) M]^{T} = M^{T}(\Sigma_{\omega e} + \Sigma_{\omega e} \bar{R}^{T} + \bar{R} \Sigma_{\omega e})
\]

\[
= M(\Sigma_{\omega e} - \Sigma_{\omega e} \bar{R} - \bar{R} \Sigma_{\omega e})
\]

(142)

since a symmetric matrix is its own transpose and a skew-symmetric matrix is the negative of its transpose.

Equations (140) and (141) are coupled with each other, and Eq. (141) is also coupled with the equations of motion of rotor and damper. Nonetheless, physically based arguments can be introduced that tend to justify the use of a transformation on \( q \), which uncouples the new deformation variables in Eq. (140) only, leaving linear combinations of these variables in each of the scalar equations of Eq. (141). Truncation of the new deformation coordinate matrix would then permit substantial reduction of the number of scalar equations accepted from Eq. (140), and this would provide exact solutions for the coordinates retained only insofar as \( \omega \) (as obtained from Eq. 141) is uninfluenced by the excluded coordinates. This seems quite an acceptable approximation, since in general the appendage vibrations act only as perturbations on the motion of the controlled vehicle, which is dominated by the control torque \( T \) in Eq. (141).

Finally, then, attention focuses on the quest for a coordinate transformation that uncouples the deformation coordinates in the 6n equations of Eq. (140). Only the homogeneous equation

\[
M' \ddot{q} + D' \dot{q} + G' \dot{q} + K' q + A' q = 0
\]

(143)

is relevant to this search.

Although matrices \( M' \) and \( D' \) are constant, matrices \( G', A', \) and \( K' \) may vary with time, since they depend on \( \omega \) and \( \dot{\omega} \). Of course \( \omega \) and \( \dot{\omega} \) are strictly unknown, but it is quite reasonable in the present context of space vehicle attitude-control-system simulation to assume that \( \omega \) varies only slightly from the nominal value (say, \( \omega(t) \)) which is the objective of the control system. Under this assumption, formal linearization removes the unknown \( \omega - \dot{\omega} \) from the homogeneous Eq. (143), since this small quantity appears in that equation only as a factor of \( \dot{q} \) or \( q \). Thus one may argue quite generally that Eq. (143) is a system of linear second-order equations with coefficients depending explicitly on time.

No general procedures are available for the transformation of Eq. (143) into a system of uncoupled equations. This can be accomplished in general terms only when the coefficient matrices are constant, which requires that the nominal value \( \omega \) be constant. This is surely the most common situation of interest in spacecraft control, but it excludes certain appendage deployment and transient operations problems.

2. State-equation modal analysis. Consider then the special case of Eq. (143) for which \( M', D', G', A' \) and \( K' \) are constant, implying \( \dot{\omega} \approx 0 \). This second-order matrix equation can always be represented by a first-order equation (the state equation). This is accomplished by defining the 12n by 1 matrix

\[
Q = \begin{bmatrix} \ddot{q} \\ \dot{q} \end{bmatrix}
\]

(144)

and writing

\[
\dot{Q} = BQ
\]

(145)

where

\[
B = \begin{bmatrix} 0 & \cdots & E \\ -M'^{-1} (K' + A') & \cdots & -M'^{-1} (G' + D') \end{bmatrix}
\]

(146)

Here \( B \) is a 12n by 12n matrix, and each of the partitions shown in Eq. (146) is of dimension 6n by 6n. Equation (145) has exactly the same content as Eq. (143), since it contains this equation in addition to the identity \( \ddot{q} = \dot{q} \).

Linear, constant-coefficient differential equations such as Eq. (145) always have solutions of the form

\[
Q = \phi^{m} e^{\lambda_{m} t}
\]

(147)

where \( \lambda_{m} \) is a scalar and \( \phi^{m} \) is a 12n by 1 matrix, which from Eq. (144) must partition into a 6n by 1 matrix \( \phi^{m} \lambda_{m} \), i.e.,

\[
\phi^{m} = [\phi_{1}^{m} \phi_{2}^{m} \cdots \phi_{6n}^{m} (\phi_{6n}^{m} \lambda_{m}) \cdots (\phi_{6n}^{m} \lambda_{m})]^{T} = -\begin{bmatrix} \phi_{1}^{m} \\ \phi_{2}^{m} \\ \vdots \\ \phi_{6n}^{m} \lambda_{m} \end{bmatrix}
\]

(148)

The validity of the solution indicated in Eq. (147) can be confirmed by substitution into Eq. (145), yielding

\[
\lambda_{m} \phi^{m} = B\phi^{m}
\]

since \( e^{\lambda_{m} t} \neq 0 \). This result may be written in the conventional form of an eigenvalue problem,

\[
(B - \lambda_{m} E) \phi^{m} = 0
\]

(149)
so that \( \lambda_m \) is an eigenvalue and \( \Phi^m \) an eigenvector of \( B \). The existence of a solution as in Eq. (147) follows from the existence of at least one eigenvalue of \( B \) and its corresponding eigenvector.

In scalar terms, Eq. (149) represents a set of \( 12n \) homogeneous algebraic equations in the \( 12n + 1 \) unknowns \( \Phi^1_m, \ldots, \Phi^n_{12n} \) and \( \lambda_m \). Nontrivial (nonzero) solutions for \( \Phi^m \) exist only when \( \lambda_m \) takes on certain characteristic values or eigenvalues such that the determinant of coefficients is zero, by Cramer's rule. In other words, values of \( \lambda_m \) must be selected that satisfy

\[
|B - \lambda_m E| = 0
\]

There are \( 12n \) solutions \( \lambda_1, \ldots, \lambda_{12n} \) for Eq. (150), as may be confirmed by expanding the determinant into the polynomial

\[
b_{12n} \lambda^{12n} + \cdots + b_1 \lambda + b_0 = (\lambda - \lambda_1)(\lambda - \lambda_2)\cdots(\lambda - \lambda_{12n}) = 0
\]

The matrix \( B \) and the related coefficients \( b_0, \ldots, b_{12n} \) are real numbers for equations that stem from the dynamic equations of Eq. (143). Therefore the complex eigenvalues among \( \lambda_1, \ldots, \lambda_{12n} \) occur in complex conjugate pairs. Subscripts may be assigned so that \( \lambda_{6n+m} = \lambda^*_m \) (asterisk here denotes complex conjugate).

Corresponding to each distinct eigenvalue \( \lambda_m \) there exists an eigenvector \( \Phi^m \) that can be determined to within a multiplicative constant by solving Eq. (149). A completely unique solution is not possible, since the system of equations in Eq. (149) includes only \( 12n - 1 \) independent algebraic equations in the \( 12n \) unknowns \( \Phi^1_m, \ldots, \Phi^n_{12n} \), with \( \lambda_m \) specified by Eq. (150). When the eigenvalues are distinct, the eigenvectors are independent (see Ref. 33, pp. 184 ff.), but when there are repeated eigenvalues, the corresponding eigenvectors may not be independent.

The validity of Eq. (149) for a particular \( \lambda_m \) and \( \Phi^m \) guarantees as well the validity of

\[
(B - \lambda_m^* E) \Phi^m = 0
\]

since the complex conjugate of zero is zero, and the conjugate of a product is the product of conjugates. Because \( \lambda_m^* = \lambda_{12n+m} \), the eigenvectors corresponding to complex pairs of eigenvalues are, from Eq. (152), also complex conjugate pairs.

Although the solution to Eq. (145) has readily been found, and certain of the properties of the solution have been noted, these steps have not yet accomplished the main objective, which is the discovery of a transformation matrix that uncouples the coordinates in Eq. (143). It is in most cases not essential that the result be a second-order differential equation; it may suffice to obtain a first-order equation such as Eq. (145), but with a diagonal coefficient matrix. Although it is possible to generate a transformation from Eq. (143) to an uncoupled system of second-order equations when \( D' \) and \( A' \) are zero, it is possible to diagonalize the state equation (145) even more generally. The latter transformation is considered first.

If \( \Phi \) is defined as the \( 12n \) by \( 12n \) matrix whose columns are the eigenvectors \( \Phi^1_m, \ldots, \Phi^{12n}_m \) of the matrix \( B \), then by inspection of Eq. (148) one may write

\[
B\Phi = B [\Phi^1 \Phi^2 \cdots \Phi^{12n}] = [B\Phi^1 \cdots B\Phi^{12n}]
\]

\[
= \begin{bmatrix} \lambda_1 & 0 & \cdots & 0 \\ 0 & \lambda_2 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & \cdots & \cdots & \lambda_{12n} \end{bmatrix}
\]

\[
= \Phi \begin{bmatrix} \lambda_1 & 0 \\ 0 & \lambda_2 \\ \vdots & \vdots \\ 0 & \lambda_{12n} \end{bmatrix}
\]

It follows that, if \( \Phi^{-1} \) exists, premultiplication provides the diagonal matrix

\[
\Phi^{-1} B\Phi = \begin{bmatrix} \lambda_1 & 0 \\ 0 & \lambda_{12n} \end{bmatrix}
\]

The existence of \( \Phi^{-1} \) is assured if the determinant of \( \Phi \) is nonzero, i.e., if the eigenvectors \( \Phi^1, \ldots, \Phi^{12n} \) are independent. It has been noted that this independence is guaranteed when \( \lambda_1, \ldots, \lambda_{12n} \) are distinct. For the dynamic system of Eq. (143), the independence of the eigenvectors can be assured even without this restriction when \( M', G', \) and \( K' \) are constant and \( A' \) and \( D' \) are zero. The necessary supporting argument is briefly summarized in the following paragraphs.

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When the eigenvectors are independent, the general solution to Eq. (145) is the linear combination of solutions of the structure of Eq. (147), i.e.,

\[ Q = \sum_{m=1}^{12n} A_m \phi_m e^{\lambda_m t} \]  
(155)

The scalar constants \( A_m \) are established by initial conditions. It follows from Eq. (155) that if the real parts of \( \lambda_1, \cdots, \lambda_{12n} \) are nonpositive, the response to initial conditions providing sufficiently small values of \( A_1, \cdots, A_{12n} \) is a solution \( Q \) with an arbitrarily small norm. In other words, when the real parts of the eigenvalues of \( B \) are nonpositive, the null solution of Eq. (145) is Liapunov stable. This is true even if there are repeated eigenvalues, as long as the eigenvectors are independent.

If on the other hand there were repeated eigenvalues (say, \( \lambda_j = \lambda_{j+1} \)) with dependent eigenvectors, the general solution would become

\[ Q = A_1 \phi^1 e^{\lambda_1 t} + \cdots + A_j \phi^j e^{\lambda_j t} + A_{j+1} \phi^{j+1} e^{\lambda_{j+1} t} + \cdots + A_{12n} \phi^{12n} e^{\lambda_{12n} t} \]  
(156)

If the real part of \( \lambda_j \) is zero, the function \( te^{\lambda_j t} \) is unbounded, and the null solution of Eq. (145) is unstable. Liapunov stability is possible in the presence of dependent eigenvectors only if the real part of the corresponding eigenvalues is negative.

Now it will be argued that the null solution of Eq. (143), restricted to constant \( M', G', K' \), and to zero values of \( A' \) and \( D' \) (and hence of the corresponding Eq. 145) is Liapunov stable, and that the eigenvalues \( \lambda_1, \cdots, \lambda_{12n} \) have nonpositive real parts. These conditions have been shown to be incompatible with the presence of dependent eigenvectors.

The function \( \dot{\mathbf{q}}^T M' \dot{\mathbf{q}} + q^T K' q \) is a Liapunov function (see Ref. 34, Ch. 1) when \( M' \) and \( K' \) are positive definite, so under these conditions the null solution of Eq. (143) (and hence of Eq. 145) is Liapunov stable. By definition (Eq. 87), \( M' \) is positive definite. The matrix \( K' \), defined by Eq. 91, consists, when \( \omega \) is constant, of the positive definite stiffness matrix \( K \) plus a matrix provided by the centrifugal "force" field induced by vehicle rotation. Unless centrifugal forces are so great as to exceed the elastic forces of the structure (in which case the appendage would pull apart), the matrix \( K' \) is positive definite. Thus under all reasonable circumstances the null solutions of Eqs. (143) and (145) are Liapunov stable, for the noted restrictions.

Because the matrix \( G' \) in Eq. (143) is skew-symmetric, and the symmetric matrix \( D' \) has been deleted, there can exist no eigenvalues with negative parts. This is a consequence mathematically of the absence of odd-powered terms in the characteristic equation (151). In physical terms, this is a reflection of the fact that the appendage as presently idealized has no damping, and consequently no attenuating solutions are admissible. Thus Liapunov stability of the null solution of Eq. (143) must mean that all eigenvalues \( \lambda_1, \cdots, \lambda_{12n} \) have zero real parts, under present restrictive assumptions.

Consequently it can be concluded that, for this class of system, the eigenvectors \( \phi^1, \cdots, \phi^{12n} \) are independent, and the matrix \( \phi^{-1} \) always exists. This means that the operation of Eq. (154) is always possible, and matrix \( B \) can always be diagonalized for the problem under consideration.

In application to the differential equations of the space vehicle, the appropriate transformation procedure is straightforward. Each of the equations involving \( q \) (Eqs. 140, 141, and 134, for example) must be rewritten in terms of \( Q \) as defined in Eq. (144). Thus Eq. (140) becomes

\[ \dot{Q} = BQ + L \]  
(157)

where, in terms of \( 6n \) by \( 1 \) submatrices \( 0 \) and \( L' \) (see Eq. 140), the \( 12n \) by \( 1 \) matrix \( L \) is \([0 \ L']^T\). Equation (141) may be written as

\[ I_0 + \omega I_0 + (i + G - h) \omega = [0_1 - (\Sigma_{ge}^T + \tilde{\Sigma} \Sigma_{ge}^T + \Sigma_{ge}^T \Sigma_{ge}) M] \dot{Q} - \dot{h} - mb \ddot{\xi} \dot{\xi}^2 + T \]  
(158)

where the \( 3 \) by \( 12n \) coefficient matrix of \( \dot{Q} \) is shown in two \( 3 \) by \( 6n \) partitions. Recall that in writing Eq. (141) from Eq. (129), the matrices \( I \) and \( G \) were noted to depend on \( q \) and \( \dot{q} \), so these matrices in Eq. (158) require reformulation in terms of \( Q \). Similarly, the damper equation (134) and any others involving \( q \) are easily rewritten in terms of \( Q \). In practice, one would normally carry the transformation into first-order equations all the way, replacing, for example, the second-order scalar damper equation by two first-order equations by introducing a \( 6 \) by \( 1 \) matrix \([\xi \ \dot{\xi}]^T\) as was done in Eq. (144), and similarly replacing \( \xi \) in Eq. (158).

The transformation

\[ Q = \phi Y \]  
(159)

is next introduced wherever \( Q \) appears in any of the equations of the vehicle or control system, and Eq. (157)
is then premultiplied by \( \Phi^{-1} \). The transformed appendage equation then becomes (by virtue of Eq. 154)

\[
\dot{\tilde{Y}} = \begin{bmatrix}
\lambda_1 & \cdots & 0 \\
0 & \ddots & 0 \\
0 & \cdots & \lambda_{12n}
\end{bmatrix} Y + \Phi^{-1} L
\]  
(160)

and the vehicle rotational equation (158) becomes

\[
I_0'' + \ddot{\omega}_0 + (i \omega + \ddot{\bar{h}}) \omega = 
[0_1 - (\sum_{E} + R_{E} + \sum_{E} r) M] \Phi \dot{\tilde{Y}} - \ddot{\bar{h}} - mb_2 E^2 + T
\]  
(161)

The scalar equations in Eq. (160) are not directly coupled by the deformation coordinates in \( Y \). It should be noted, however, that the matrix \( L \) in Eq. (160) is determined by the base rotational motion obtained from Eq. (161), and in general, all of the scalar variables in \( Y \) participate in Eq. (161). Thus when the total system of differential equations is considered, there is still coupling among the deformation coordinates in \( Y \). Truncation of the \( 12n \) by 1 matrix \( Y \) and the corresponding \( 12n \) by \( 12n \) transformation matrix \( \Phi \) must therefore be based on the considered judgment that the coordinate coupling is weak and physically indirect. Such arguments support the practice of determining coordinate truncation principally from the uncoupled appendage deformation equations in Eq. (160). It may be necessary to consider also certain characteristics of the control system, such as the range of frequencies of response, sensor data frequency filtering, and the location of sensors and thrustors. Discussion of these questions of control system interaction is postponed until Section V.

The homogeneous solution for \( Q \) in Eq. (155) is, of course, the same as that obtained by solving Eq. (160) for \( Y \) when \( L \) is zero and substituting into the transformation of Eq. (159). Truncation of \( Y \) is normally accomplished by inspection of this homogeneous solution, although it may also be necessary to include any coordinates in \( Y \) that may introduce resonances due to coincidence of the imaginary part of the corresponding \( \lambda \) with a driving frequency in the forcing term \( L \) in Eq. (160). Since truncation is determined from physical considerations, it is important that the physical significance of the variables \( Y_1, \ldots, Y_{12n} \) be understood. This may seem difficult, since complex eigenvectors in \( \Phi \) make the coordinates \( \tilde{Y}_1, \ldots, \tilde{Y}_{12n} \) generally complex, even when \( Q \) is by definition real (see Eq. 159). The indicated homogeneous solution of Eq. (160) has the elements

\[
Y_m = A_m e^{\lambda_m t}
\]  
(162)

Thus the solution shown in Eq. (155) may be written as

\[
Q = \sum_{m=1}^{12n} Y_m \phi_m
\]  
(163)

in conformance with Eq. (159). Although individual scalars \( Y_m \) and column matrices \( \phi_m \) are complex, the reality of \( Q \) is assured by the appearance of eigenvalues and eigenvectors in complex pairs. Consider, for example, a solution for \( Y \) with initial conditions selected so that all values of \( A_m \) in Eq. (162) are zero except \( A_\kappa \) and \( A_{\kappa+\kappa} \). Then \( Q \) is

\[
Q = Y_\kappa \phi_\kappa + Y_{\kappa+\kappa} \phi^{\kappa+\kappa}
\]

\[
= A_\kappa \phi_\kappa e^{\lambda_\kappa t} + A_{\kappa+\kappa} \phi^{\kappa+\kappa} e^{\lambda_{\kappa+\kappa} t}
\]

\[
= A_\kappa \phi_\kappa e^{\lambda_\kappa t} + A_{\kappa+\kappa} \phi^{\kappa+\kappa} e^{\lambda_{\kappa+\kappa} t}
\]

Let \( \lambda_\kappa = \alpha_\kappa + i\omega_\kappa \) and \( \phi_\kappa = \psi_\kappa + i\Gamma_\kappa \) and substitute \( e^{\lambda_\kappa t} = e^{\alpha_\kappa t} (\cos \omega_\kappa t + i \sin \omega_\kappa t) \), etc. Then \( Q \) becomes

\[
Q = e^{\alpha_\kappa t} \left( (A_\kappa + A_{\kappa+\kappa}) (\psi_\kappa \cos \omega_\kappa t - \Gamma_\kappa \sin \omega_\kappa t) \right)
\]

\[
+ i(A_\kappa - A_{\kappa+\kappa}) (\Gamma_\kappa \cos \omega_\kappa t + \psi_\kappa \sin \omega_\kappa t)
\]

Since \( Q \) is real, \( A_{\kappa+\kappa} \) is \( A^*_\kappa \). Let \( A_\kappa = (C_\kappa - iD_\kappa)/2 \), so that \( Q \) may be rewritten in wholly real terms as

\[
Q = e^{\alpha_\kappa t} \left( C_\kappa (\psi_\kappa \cos \omega_\kappa t - \Gamma_\kappa \sin \omega_\kappa t) \right)
\]

\[
+ D_\kappa (\psi_\kappa \sin \omega_\kappa t + \Gamma_\kappa \cos \omega_\kappa t)
\]

(164)

Thus the general homogeneous solution for \( Q \) appears in real terms as

\[
Q = \sum_{m=1}^{6n} e^{\alpha_m t} \left( C_m (\psi_m \cos \sigma_m t - \Gamma_m \sin \sigma_m t) \right) + D_m (\psi_m \sin \sigma_m t + \Gamma_m \cos \sigma_m t)
\]  
(165)

The \( 12n \) scalars \( C_1, \ldots, C_{6n} \) and \( D_1, \ldots, D_{6n} \) are established by the initial conditions on \( Q \). For an undamped system, as in Eq. (143) when \( D' = 0 \), the scalars \( \alpha_1, \ldots, \alpha_{6n} \) are all zero.

Now coordinate truncation can be based on the same rationale traditionally employed by structural dynamicists in application to second-order equations of motion. Each conjugate pair of coordinates (say, \( Y_m \) and \( Y_{\kappa+\kappa} = Y^*_m \)) corresponds to appendage vibrations of the entire structure at a given frequency (here \( \sigma_m \)). In general, these vibrations are not in phase throughout the structure, as would be the case for classic normal modes of vibration about a rest configuration. The classic interpretation of the eigenvector \( \phi^*_m \) of Eq. (148) as the mode shape of the...
vibration at frequency \( \sigma_m \) is modified by the complex nature of \( \Phi^m = \Phi^m + i \Phi^m \). As illustrated in real terms in Eq. (164), this is reflected as two different mode shapes, corresponding to two vibrations at frequency \( \sigma_m \), with a 90-deg phase lag. Furthermore, these mode shapes depend on the initial conditions. The coordinates in \( Y \) are still called modal coordinates in this report, despite these deviations from the classic interpretation.

The usual decision of the structural dynamicist is to retain in truncation the responses at the lowest natural frequencies, and also any others at which expected driving frequencies are close to natural frequencies (with consequent danger of resonance). The justification for concentrating on lower frequency responses is often based on the fact that storing a given amount of energy as potential energy or strain energy in the deformation of an elastic structure into a low-frequency mode shape generally results in larger displacements and base reactions than are present when the same energy is stored in a high-frequency mode shape deformation. To this it may be added that the present elastic model is unrealistic in its neglect of damping, and that, in fact, the higher frequency vibrations will have more damping than those at low frequency. This again supports the practice of ignoring the high-frequency response (since it rapidly attenuates in the transient solution anyway).

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The appendage deformation matrix \( Y \) in Eqs. (160) and (161) is therefore replaced by the \( 2N \) by 1 truncated matrix

\[
\bar{Y} = [Y_1 \cdots Y_N Y_1^* \cdots Y_N^*]^T \quad (166)
\]

where \( N \) is the number of modes to be preserved in the simulation. The transformation matrix \( \Phi \) is accordingly truncated to the \( 12n \) by \( 2N \) matrix \( \bar{\Phi} \), where

\[
\bar{\Phi} = [\phi^1 \cdots \phi^N \phi^{1*} \cdots \phi^{N*}] \quad (167)
\]

The equation of motion of the appendage (Eq. 160) now becomes

\[
\dot{\bar{Y}} = \begin{bmatrix} \lambda_1 & 0 \\ \lambda_N^* & \lambda_N^* \\ 0 & \lambda_N^* \end{bmatrix} \bar{Y} + \bar{\Phi}^{-1} L \quad (168)
\]

This matrix equation consists of \( 2N \) scalar equations in the complex variables \( Y_1, \cdots, Y_N, Y_1^*, \cdots, Y_N^* \). In computation, only the first \( N \) equations need be considered, since the equations are uncoupled and the second set of \( N \) variables is available as the complex conjugates of the first set.

The rotational equation of the vehicle (Eq. 161) then becomes

\[
I_\omega + \omega \omega + (I + G - \bar{h}) \omega = \left[ 0_1 - (\Sigma_{\phi_i} + \bar{R}_S \Sigma_{\phi_i} + \Sigma_{\phi_i} m) \right] \bar{\Phi} \bar{Y} - \bar{h} - mb \bar{E}^2 + T
\]

(169)

The matrix \( \bar{\phi} \) is rectangular, and the symbol \( \bar{\phi}^{-1} \) in Eq. (168) represents a left inverse of \( \bar{\phi} \). This is available (Ref. 35) as

\[
\bar{\phi}^{-1} = (\bar{\phi}^T \bar{\phi})^{-1} \bar{\phi}^T
\]

(170)

as may be confirmed by the multiplication

\[
[(\bar{\phi}^T \bar{\phi})^{-1} \bar{\phi}^T] \bar{\phi} = (\bar{\phi}^T \bar{\phi})^{-1} (\bar{\phi}^T \bar{\phi}) = E
\]

The inversion actually performed in Eq. (170) to obtain \( \bar{\phi}^{-1} \) is of a matrix of dimension \( 2N \). Computations are thus not as imposing a task as may have been implied by the presence of the \( 12n \) by \( 12n \) matrix inverse \( \bar{\phi}^{-1} \) in Eq. (160). In practice, the numerical value of \( \bar{\phi} \) is never computed, since only \( \bar{\phi} \) is required.

Incorporation of energy dissipation into the appendage mathematical model is usually deferred, pending transformation to modal coordinates. The scalar \( \sigma_m \) in Eq. (164) would be zero for the undamped system of Eq. (143) with \( D' = 0 \), since \( G' \) is skew-symmetric. Conventional practice in structural dynamics is to assign values to \( \sigma_1, \cdots, \sigma_N \) (where \( N \) modes are retained in the modal-coordinate truncation), with preliminary numerical values based on experience with similar structures and final values based on prototype tests. This amounts to rather arbitrarily adding a viscous damping term \( D' \dot{q} \) to Eq. (143) after completing the modal analysis, and assuming that the structure of the matrix \( D' \) is such that the eigenvectors \( \phi^1, \cdots, \phi^{12n} \) are undisturbed by this addition. Numerical values of the elements in \( D' \) are never considered, since they are determined implicitly by the choice of \( \sigma_1, \cdots, \sigma_N \). (Actually, the conventional pattern is the assignment of a value to \( \xi_m = \sigma_m / \sigma_n \) rather than to \( \sigma_n \) itself. Here \( \xi_m \) is called the “percentage of critical damping.” For typical spacecraft appendage structures, \( \xi_m \) is well below 5% for the lower frequency responses.)
The mechanical process of incorporating damping in the appendage equations is straightforward. Equation (168) includes a diagonal matrix of eigenvalues \( \lambda_1, \ldots, \lambda_N \), \( \lambda_1^* \cdot \ldots \cdot \lambda_N^* \). As calculated from the matrix \( B \) of Eq. (146) with \( D' \) set equal to zero, these eigenvalues will be purely imaginary, e.g., \( \lambda_m = i \sigma_m \). To incorporate structural damping, one instead substitutes \( \lambda_m = \xi_m \sigma_m + i \sigma_m \) into Eq. (168), where \( \xi_m \) is chosen as noted in the preceding paragraph.

It should be noted that the procedure just described for transformation to modal coordinates and subsequent truncation did not require any special properties of the constant matrix \( B \) in Eq. (145), except in proving the existence of \( \Phi^{-1} \) in Eq. (154). The nonsingularity of \( \Phi \) was established by relying upon the skew-symmetry of matrix \( G' \) and the absence of the viscous damping matrix \( D' \) in Eq. (143). This property was used to prove that even in the special case of repeated eigenvalues, the independence of the eigenvectors is assured, so the solution to Eq. (145) has the form of Eq. (155) rather than that of Eq. (156). The proof would proceed equally well if \( G' \) were zero, but when damping is present the formal argument underlying the previous discussion breaks down. Although it seems reasonable that continuity would require the preservation of the nonsingularity of \( \Phi \) in the presence of small damping forces (with the corresponding \( D' \hat{q} \) included in Eq. 143, with \( D' \) symmetric), proof of this condition seems lacking.

It can be asserted, then, that only when \( \Phi^{-1} \) exists is the procedure applied here to transform the state equation (145) into the diagonal form applicable. It is perhaps worth repeating that the existence of \( \Phi^{-1} \) is assured when the eigenvalues of \( B \) are distinct, or when the matrix \( D' \) is zero. This method has thus been shown to be applicable to any linear dynamic system with distinct natural frequencies, and to any undamped linear dynamic system. Applicability to damped linear dynamic systems with repeated natural frequencies seems probable, but has not been formally established.

### 3. Modal analysis of damped, nonrotating structures.

The utility of the direct analysis of the state equation may depend on the computational efficiency of the inversion of \( \Phi \) (after truncation). It should be noted that \( \Phi \) is not orthogonal, i.e., \( \Phi^{-1} \neq \Phi^T \). Foss (Ref. 36) has shown that when \( G' = A' = 0, D' \) is nonzero, and eigenvalues are distinct, certain weighted orthogonality relationships can be established. Foss notes the equivalence of Eqs. (157) and (140) with the first-order equation

\[
M'' \dot{\Phi} + K'' \Phi = L''
\]

where

\[
M'' = \begin{bmatrix} D' & M' \\ M' & 0 \end{bmatrix}
\]
\[
K'' = \begin{bmatrix} K' & 0 \\ 0 & -M' \end{bmatrix}
\]
\[
L'' = \begin{bmatrix} L' \\ 0 \end{bmatrix}
\]

Foss demonstrates that, for symmetric \( D' \), the eigenvectors \( \Phi^m \) and \( \Phi^k \) corresponding to different eigenvalues \( (\lambda_m \neq \lambda_k) \) have the orthogonality property

\[
\Phi^m^T M'' \Phi^k = 0
\]
\[
\Phi^m^T K'' \Phi^k = 0
\]

Thus for suitable eigenvector normalization, diagonalization of Eq. (171) with the transformation (see Eq. 159)

\[
\begin{bmatrix} q \\ \dot{q} \end{bmatrix} \equiv \Phi Y
\]

is computationally easier than the diagonalization of Eq. (145). Whereas the latter procedure requires the calculation of \( \Phi^{-1} \) (or \( \Phi^{-1} \) after truncation), the former requires only the construction of \( \Phi^T \) (or \( \Phi^T \)) to obtain the same result. This transformation of Eq. (171), followed by premultiplication by \( \Phi^T \), provides the same results as Eq. (168), namely,

\[
\Phi^T M'' \Phi Y + \Phi^T K'' \Phi Y = \Phi^T L''
\]

or, with Eq. (172) and truncation,

\[
\begin{bmatrix} \lambda_1 & \ldots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \ldots & \lambda_N \end{bmatrix} \begin{bmatrix} \lambda_1^* & \ldots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \ldots & \lambda_N^* \end{bmatrix} \bar{Y} = \bar{\Phi}^T L''
\]

\[\text{JPL TECHNICAL REPORT 32-1329}\]
This equation combines with the vehicle equation (169) and parallel equations for rotor and damper to obtain a complete simulation. Equation (173) is not applicable to the differential equations of an appendage on a rotating base (Eqs. 140 and 143 with \( G' \neq 0 \)). When the base is not rotating, and specific damping forces are necessary (perhaps because of discrete dashpots in the system), the method of Foss is computationally superior, but mathematically equivalent, to the more general method used to derive Eq. (168). Both of these methods have two major undesirable features:

(1) Except in the very special case for which \( G' = 0 = A' \) and \( D' \) is a polynomial in the symmetric matrices \( M' \) and \( K' \) (Ref. 30), the eigenvectors forming the columns of the transformation matrix \( \Phi \) are complex, and thus the new deformation variables \( Y \) introduced in Eq. (159) are also complex. Although physical interpretation of the complex coordinates (as conjugate pairs) has been provided, this interpretation is less immediate and probably less comfortable for the engineer than would be a system of real coordinates. New computer programs might be necessitated by the simulation in terms of complex coordinates, whereas existing programs might suffice if a useful transformation to real coordinates can be found.

(2) The final equation of motion of the appendage (Eq. 168 or Eq. 173) is a first order matrix equation. Although controls engineers are finding increasing use for first-order equations, the structural dynamicist is more accustomed to working with second-order equations. Computation with existing computer programs may be precluded and interpretation of results may be impeded by the adoption of first-order equations.

4. Modal analysis of undamped, rotating structures.* When in Eq. (143) the acceleration matrix \( A' \) and the damping matrix \( D' \) are zero, an alternative to the transformation of Eq. (159) can be devised that results ultimately in a system of uncoupled, real, second-order scalar equations of motion. This method is inapplicable when damping is included in the mathematical model of the appendage structure, although the introduction of modal damping into the transformed equations is not precluded. The transformation to be described will uncouple the homogeneous second-order appendage equations even when the appendage is on a rotating base (provided that \( A' = 0 \) in Eq. 143), but this will not be successful if a discrete dashpot is incorporated into the appendage or the appendage support structure, since then \( D' \neq 0 \) in Eq. (143).

The general solution for Eq. (143) with constant coefficients has already been found, since \( q \) is identified in Eq. (144) as the upper half of \( Q \), and the solution for \( Q \) has been recorded in real terms in Eq. (165). As noted previously, \( \sigma_m \) is zero for all \( m \) for undamped systems. If now the upper halves of \( \psi \) and \( \Gamma \) are denoted by the corresponding lower-case letters (as in Eq. 148 for \( \phi \)), the substitutions

\[
\begin{bmatrix} \phi^m \\ \gamma^m \end{bmatrix} = 
\begin{bmatrix} \psi^m \\ \gamma^m \lambda_m \end{bmatrix}
\]

(174)

permit \( q \) to be written from Eq. (165) in the form

\[
q = \sum_{m=1}^{en} \left\{ C_m (\psi^m \cos \sigma_m t - \gamma^m \sin \sigma_m t) + D_m (\psi^m \sin \sigma_m t + \gamma^m \cos \sigma_m t) \right\}
\]

(175)

The derivative \( \dot{q} \) may be obtained by differentiating \( q \) or by inspecting the bottom half of Eq. (165), as

\[
\dot{q} = \sum_{m=1}^{en} \left\{ -C_m \sigma_m (\psi^m \sin \sigma_m t + \gamma^m \cos \sigma_m t) + D_m \sigma_m (\psi^m \cos \sigma_m t - \gamma^m \sin \sigma_m t) \right\}
\]

(176)

If now \( Z^h \) is defined as the \( 12n \) by 1 matrix

\[
Z^h =
\begin{bmatrix}
C_1 \cos \sigma_1 t + D_1 \sin \sigma_1 t \\
C_2 \cos \sigma_2 t + D_2 \sin \sigma_2 t \\
\vdots \\
C_{en} \cos \sigma_{en} t + D_{en} \sin \sigma_{en} t \\
-C_1 \sin \sigma_1 t + D_1 \cos \sigma_1 t \\
\vdots \\
-C_{en} \sin \sigma_{en} t + D_{en} \cos \sigma_{en} t
\end{bmatrix}
\]

(177)

then by inspection of Eqs. (165) and (177), one may write

\[
Q = PZ^h
\]

(178)

*Require constant nominal rotation rate, and \( A' = 0 \). See footnote on page 28.
where \( P \) is the transformation matrix

\[
P = \begin{bmatrix}
\varphi^1 & \varphi^2 & \cdots & \varphi^{6n} \\
-\sigma_1 \varphi^1 & -\sigma_2 \varphi^2 & \cdots & -\sigma_{6n} \varphi^{6n} \\
\vdots & \vdots & \ddots & \vdots \\
-\sigma_{6n} \varphi^{6n} & -\sigma_{6n} \varphi^{6n} & \cdots & -\sigma_{6n} \varphi^{6n}
\end{bmatrix}
\] (179)

By inspection of the matrix \( Z^h \) and its time derivative, one may write

\[
\dot{Z}^h = \begin{bmatrix}
0 & \sigma_1 & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & -\sigma_{6n} & -\sigma_{6n} \\
\end{bmatrix}
\]

\[
Z^h = \begin{bmatrix}
0 & \sigma \\
-\sigma & 0 \\
\vdots & \vdots \\
0 & -\sigma_{6n}
\end{bmatrix}
\] (180)

where \( \sigma \) has been defined as the \( 6n \) by \( 6n \) diagonal matrix of natural frequencies \( \sigma_1, \cdots, \sigma_{6n} \).

The first \( 6n \) elements of \( Z^h \) may be assigned the symbols \( z_1, \cdots, z_{6n} \), so that Eq. (177) may be written as

\[
Z^h = \begin{bmatrix}
z_1 \\
\vdots \\
z_{6n}
\end{bmatrix} = \begin{bmatrix}
\sigma_1 z_1 \\
\vdots \\
\sigma_{6n} z_{6n}
\end{bmatrix}
\] (181)

which defines the \( 6n \) by 1 column matrix \( z \) and the \( 6n \) by \( 6n \) diagonal matrix \( \sigma^{-1} \) whose elements are the frequency reciprocals \( 1/\sigma_1, \cdots, 1/\sigma_{6n} \). In terms of \( z_1, \cdots, z_{6n} \), Eq. (180) becomes the set of scalar equations

\[
\begin{align*}
\dot{z}_1 &= \dot{z}_1 \\
\vdots \\
\dot{z}_{6n} &= \dot{z}_{6n} \\
\ddot{z}_1 &= -\sigma_1 \dot{z}_1 \\
\vdots \\
\ddot{z}_{6n} &= -\sigma_{6n} \dot{z}_{6n}
\end{align*}
\] (182)

The first set of \( 6n \) scalar equations in Eq. (182) is trivial, but the second set is a group of \( 6n \) uncoupled equations in the real variables \( z_1, \cdots, z_{6n} \), which is equivalent to Eq. (143). In matrix terms, this result is

\[
\ddot{z} + \sigma^2 z = 0
\] (183)

where

\[
\sigma^2 = \begin{bmatrix}
\sigma_1^2 & 0 \\
0 & \sigma_{6n}^2
\end{bmatrix}
\]

To accomplish a direct transformation from \( q \) in Eq. (143) to \( z \) in Eq. (183), rewrite the transformation in Eq. (144) as

\[
q = [E_1 \ 0] Q
\] (184)

where the identity matrix \( E \) and the null matrix \( 0 \) are of dimension \( 6n \) by \( 6n \). Substitution of Eqs. (178) and (181) into Eq. (184) produces

\[
q = \begin{bmatrix}
\zeta \\
\ddot{\zeta}
\end{bmatrix}
\] (185)

or, from Eq. (179),

\[
q = \psi \dot{z} + \gamma \sigma^{-1} \ddot{z}
\] (186)

where \( \psi \) and \( \gamma \) are the \( 6n \) by \( 6n \) matrices

\[
\psi = \begin{bmatrix}
\varphi^1 \varphi^2 & \cdots & \varphi^{6n}
\end{bmatrix}
\]

and

\[
\gamma = \begin{bmatrix}
\gamma^1 \gamma^2 & \cdots & \gamma^{6n}
\end{bmatrix}
\]

A transformation of the character of Eq. (185), which relates old coordinates to a combination of new coordinates and their derivatives (or, more often, their generalized momenta), is known as a contact transformation. In contrast, a transformation that relates one set of generalized coordinates directly to another is called a point transformation. As noted in Whittaker (Ref. 24, p. 195, footnote), there exists no point transformation that takes equations of the structure of Eq. (143) with \( A' = D' = 0 \).
into uncoupled second-order equations such as Eq. (183), unless $G'$ is also zero. The existence of a contact transformation that accomplishes this objective when $G' \neq 0$ is noted in Whittaker (p. 427), where equations of motion are expressed in terms of Hamilton’s canonical equations.

Although Eq. (143) has been the starting point in the quest for useful transformations, it is the inhomogeneous counterpart, Eq. (140), that must actually be employed in the simulation. In terms of first-order equations, it is Eq. (157) and not Eq. (145) that must be used. Direct substitution of the transformation in Eq. (186) into the second-order inhomogeneous Eq. (140) does not appear to be fruitful. Instead the transformation

$$Q = PZ$$

is used in the first-order inhomogeneous Eq. (157), where $P$ is a matrix of constants available from the eigenvalues and eigenvectors of $B$ as in Eq. (179), and $Z$ is a new matrix of unknowns. The result is, from Eqs. (157) and (187),

$$P\dot{Z} = BPZ + L$$

Only when $L$ is zero is the solution for $Z$ that given as $Z \dot{h}$ by Eq. (177).

When $P$ is nonsingular (a condition not always assured), Eq. (188) may be written as

$$\dot{Z} = P^{-1}BPZ + P^{-1}L$$

The matrix $P^{-1}BP$ has been shown in the context of the analysis of the homogeneous equation (see Eq. 180) to have the value

$$P^{-1}BP = \begin{bmatrix} 0 & 0 \\ -\sigma & 0 \end{bmatrix}$$

and this product of constant matrices cannot change its value when the equation becomes inhomogeneous.

The inverse of $P$ as required in Eq. (189) is expressed and computed most easily in terms of its partitioned elements, according to the formula (Ref. 37, p. 640)

$$P^{-1} = \begin{bmatrix} (P_{11} - P_{12}P_{21}P_{22})^{-1} & -P_{11}P_{12}(P_{22} - P_{21}P_{11}P_{12})^{-1} \\ -P_{22}P_{21}(P_{11} - P_{12}P_{21}P_{22})^{-1} & (P_{22} - P_{21}P_{11}P_{12})^{-1} \end{bmatrix}$$

(191)

where

$$P = \begin{bmatrix} P_{11} & P_{12} \\ P_{21} & P_{22} \end{bmatrix} = \begin{bmatrix} \psi & \gamma \\ -\sigma \gamma & \sigma \psi \end{bmatrix}$$

(192)

With the identity typified by

$$(\sigma \psi)^{-1} = \psi^{-1} \sigma^{-1}$$

(193)

the inverse of $P$ simplifies to

$$P^{-1} = \begin{bmatrix} (\psi + \gamma \psi^{-1} \gamma)^{-1} & -\psi^{-1} \gamma (\psi + \gamma \psi^{-1} \gamma)^{-1} \sigma^{-1} \\ \psi^{-1} \gamma (\psi + \gamma \psi^{-1} \gamma)^{-1} & (\psi + \gamma \psi^{-1} \gamma)^{-1} \sigma^{-1} \end{bmatrix}$$

(194)
In expanded form, Eq. (189) may now be written (see Eq. 157 for $L$ expansion) in the form

$$
\begin{bmatrix}
Z_1 \\
\vdots \\
Z_{12n}
\end{bmatrix}
= \begin{bmatrix}
0 & \sigma_1 & 0 \\
& \ddots & \ddots \\
& & 0 & \sigma_{6n}
\end{bmatrix}
\begin{bmatrix}
Z_1 \\
\vdots \\
Z_{12n}
\end{bmatrix}
$$

$$
+ \begin{bmatrix}
(\psi + \gamma \psi^{-1} \gamma)^{-1} \\
\psi^{-1} (\psi + \gamma \psi^{-1} \gamma)^{-1}
\end{bmatrix}
\begin{bmatrix}
-\psi^{-1} \gamma (\psi + \gamma \psi^{-1} \gamma)^{-1} \sigma_{6n} \\
(\psi + \gamma \psi^{-1} \gamma)^{-1} \gamma
\end{bmatrix}
\begin{bmatrix}
0 \\
L'
\end{bmatrix}
$$

(195)

In the analysis of the homogeneous equation, with $L'$ absent, the elements of the column matrix $Z$, labeled $Z'$, can be identified as in Eq. (181). This permits the scalar equations obtained from the upper half of $Z$ to be discarded as trivial identities (see Eq. 182), and leaves the lower half of $Z$ to provide nontrivial, uncoupled second-order equations in the new variables $z_1, \ldots, z_{6n}$ (Eq. 183). This identification and resolution into second-order equations does not appear to be possible in the inhomogeneous case (Eq. 195), unless $-\psi^{-1} \gamma (\psi + \gamma \psi^{-1} \gamma)^{-1} \sigma_{6n}$ is zero. Note that the inverted matrix in parentheses cannot be zero, or every term in $P^{-1}$ is zero, and the applied forces and torques in $L'$ have no dynamic consequences. The most useful condition under which Eq. (195) permits this identification of the elements of $Z$ as in Eq. (181) is the condition $\gamma = 0$. This corresponds to the special case of real eigenvectors, which implies the absence of $G'$ in Eqs. (143) and (140). This is then the classic problem of the vibration of a structure about a state of inertial rest, to which attention will return shortly.

In general, the $12n$ first-order equations in Eq. (195) do not admit of reduction to second-order equations in $z$ as in the homogeneous case (Eq. 183). This reduction does serve, nonetheless, to motivate the coordinate truncation of the deformation variable matrix $Z$ in the inhomogeneous Eq. (195).

With the use of frequency criteria, the truncation of the $6n$ by $1$ matrix $z$ in Eq. (183) is completely straightforward; one simply ignores all but $z_1, \ldots, z_N$ of the elements in $z$ when frequencies $\sigma_1, \ldots, \sigma_{6n}$ are of interest. The coordinates $z_1, \ldots, z_N$ are modal coordinates in the broad sense in which that term is used in this report. If the solutions to the scalar equations in Eq. (183) are written as

$$
z_m = C_m \cos \sigma_m t + D_m \sin \sigma_m t \quad \text{or} \quad z_m = -C_m \sigma_m^{-1} \cos \sigma_m t + D_m \sigma_m^{-1} \sin \sigma_m t$$

then substitution into Eq. (186) yields the solution for $q$ in Eq. (175), for which physical interpretation has been provided.

The introduction of energy-dissipation capability into the mathematical model can also be accomplished in the homogeneous case after transformation to the modal-coordinate matrix $z$. Equation (183) is simply replaced by

$$\ddot{\zeta} + 2\zeta_0 \dot{\zeta} + \zeta_0^2 \zeta = 0$$

(197)

where the overbars indicate truncation to $N$ coordinates, and

$$\zeta = \begin{bmatrix}
z_1 \\
\vdots \\
z_N
\end{bmatrix}$$

In the inhomogeneous case, which is of primary interest, the matrix $Z$ in Eq. (195) can be written in partitioned terms as

$$Z = \begin{bmatrix}
Z' \\
Z^2
\end{bmatrix}$$

(198)

where $Z'$ and $Z^2$ are $6n$ by $1$ matrices. The transformation derived as Eq. (186) for the homogeneous case then becomes

$$q = \psi Z' + \gamma Z^2$$

(199)
While it is no longer true in general that \( Z_1^m \) and \( Z_2^m \) are related by \( Z_1^m = \sigma_m Z_2^m \) as was the case for the homogeneous solution, there remains an association between these two coordinates and the frequency \( \sigma_m \). If the corresponding scalar equations are isolated in Eq. (185), they appear as

\[
\dot{Z}_1^m = \sigma_m Z_2^m - [\psi^{-1} \gamma (\psi + \gamma^{-1} \gamma^{-1})^{-1} \sigma^{-1}]_m L'
\]

and

\[
\dot{Z}_2^m = -\sigma_m Z_1^m + [\psi + \gamma^{-1} \gamma^{-1}]_m L'
\]

where, when \( N \) modes are retained,

\[
\tilde{Z}_1 = \begin{bmatrix} Z_1^1 \\ \vdots \\ Z_1^N \end{bmatrix}, \quad \tilde{Z}_2 = \begin{bmatrix} Z_2^1 \\ \vdots \\ Z_2^N \end{bmatrix}
\]

\[
\tilde{\sigma} = \begin{bmatrix} \sigma & 0 \\ \vdots & \vdots \\ 0 & \sigma_X \end{bmatrix}, \quad \tilde{\psi} = [\psi^1 \psi^2 \cdots \psi^N]
\]

\[
\tilde{\gamma} = [\gamma^1 \gamma^2 \cdots \gamma^N]
\]

Modal damping may be introduced into these equations in such a way that they reduce in the homogeneous case to Eq. (197). This can be accomplished by inserting the matrix \(-2 \tilde{\zeta} \tilde{\sigma}\) (see Eq. 197) at the appropriate place in Eq. (200). The result is recorded as Eq. (201), which employs the symbol \( \tilde{P}^{-1} \) for the truncated \( P \) inverse written in more detail in Eq. (200):

\[
\begin{bmatrix} \dot{\tilde{Z}}_1 \\ \dot{\tilde{Z}}_2 \end{bmatrix} = \begin{bmatrix} 0 & \tilde{\sigma} \\ \tilde{\sigma} & -2 \tilde{\zeta} \tilde{\sigma} \end{bmatrix} \begin{bmatrix} \tilde{Z}_1 \\ \tilde{Z}_2 \end{bmatrix} + \tilde{P}^{-1} \begin{bmatrix} 0 \\ L' \end{bmatrix}
\]

This equation is in final form for the simulation of a flexible appendage on a controlled vehicle. It is therefore an alternative to Eq. (168). Each of the three appendage equations (168), (173), and (201), has advantages and disadvantages.

5. Comparison of three alternative transformations. The three coordinate transformations considered thus far do not exhaust the useful possibilities; indeed, the most commonly used transformations have yet to be considered. It may nonetheless be appropriate to pause to review and summarize these three transformations.

The most general form of the appendage equation is Eq. (168), since it accommodates a discretely damped appendage on a base rotating at a constant rate. In other words, it is applicable to a discrete-coordinate appendage equation of the form

\[
M' \ddot{q} + D' \dot{q} + G' q + K' q + A' q = L'
\]

where the subscript \( m \) outside of the square brackets denotes the \( m \)th row of the matrix within the brackets. Again it is evident that if either \( \gamma \) or \( L' \) is zero, then \( Z_1^m = Z_2^m/\sigma_m \). Even in the general case, however, it is most reasonable to truncate the matrices \( Z_1^m \) and \( Z_2^m \) by retaining only those elements \( Z_{1m}^m \) and \( Z_{2m}^m \) corresponding to frequencies \( \sigma_m \) that are of interest. The matrices of real and imaginary parts of eigenvectors, \( \psi \) and \( \gamma \), are truncated correspondingly. Truncated matrices are denoted by an overbar, as previously, so Eq. (195) appears after truncation as

\[
\tilde{Z}_1 = [\psi^{-1} \gamma (\psi + \gamma^{-1} \gamma^{-1})^{-1} \tilde{\sigma}]_m L'
\]

\[
\tilde{Z}_2 = [-\tilde{\sigma} \tilde{\psi}^{-1} \gamma (\psi + \gamma^{-1} \gamma^{-1})^{-1} \tilde{\sigma}]_m L'
\]
corresponding discrete coordinate equations are

\[ M' \ddot{q} + D' \dot{q} + K' q = L' \]  

(203)

with all coefficient matrices symmetric and positive definite. If \( D' \) is not a polynomial in \( M' \) and \( K' \) (see Ref. 30), the method of Foss (Eq. 173) is most advantageous, for the reasons noted in the text preceding Eq. (173).

For an undamped elastic appendage on a rotating base, * Eq. (140) is the discrete-coordinate equation

\[ M' \ddot{q} + G' \dot{q} + K' q = L' \]  

(204)

This equation has been shown to be equivalent to Eq. (200), which has the advantage over Eq. (168) of involving only real numbers (recall that in Eq. 168 the coordinates in \( \Phi \) are complex, as are the eigenvectors in \( \Phi \), which must be inverted).

Equation (200) was modified and written as Eq. (201), which includes modal damping. This step cannot be justified in any formal way, since it obviously involves a change in the mathematical model of the physical system. Modal damping is introduced in Eq. (201) in such a way that the homogeneous equation reduces to Eq. (197), which yields a solution corresponding to Eq. (165). This is at best a crude attempt to provide some mathematical representation of the oscillation attenuation that must occur for a real structure. In fact, the substitution of the homogeneous equation reduces to Eq. (197), which is of the form of Eq. (139). One can see that this equation provides a simulation of an appendage attached to a base that is nominally not rotating in inertial space. If the base angular-velocity matrix \( \omega \) in Eq. (139) is assumed to remain small, and higher degree terms in \( \omega \), \( \dot{\omega} \), and \( \ddot{\omega} \) are ignored, then \( \omega \) may be replaced by \( h \), and Eq. (139) becomes

\[ M' \ddot{q} + K' q = L' \]  

(206)

which is at once the simplest and most useful system of equations. By returning to the appendage equation in its explicit form (Eq. 139), one can see that this equation provides a simulation of an appendage attached to a base that is nominally not rotating in inertial space. If the base angular-velocity matrix \( \omega \) in Eq. (139) is assumed to remain small, and higher degree terms in \( \omega \), \( \dot{\omega} \), and \( \ddot{\omega} \) are ignored, then \( \omega \) may be replaced by \( h \), and Eq. (139) becomes

\[ M (E - \Sigma_{g=0}^{s} \Sigma_{e=0}^{r} \frac{M}{\Omega^2} \xi) \ddot{q} + Kq = \]

\[ -M (\Sigma_{g=0}^{s} \Sigma_{e=0}^{r} \xi) \ddot{q} \]

\[ -M \Sigma_{g=0}^{s} F / \Omega + M \Sigma_{g=0}^{s} m \ddot{E} / \Omega + \lambda \]  

(207)

which is of the form of Eq. (206).

This simplest case is of course amenable to analysis by any of the three methods discussed earlier, but more efficient procedures can be found in any textbook on structural dynamics (e.g., Ref. 11). Because of the availability of proofs in standard references, the classic techniques of the structural dynamicist are applied here to Eqs. (206) and (207) without proofs.

As noted in a somewhat broader context in Eq. (147), the homogeneous solution to Eq. (206), or (207), has the form

\[ q = \sum_{j=1}^{n} c_j e^{\lambda_j t} \phi_j \]  

(208)
where $\lambda_j$ and $\phi_j$ are eigenvalues and eigenvectors available from

$$[M' \lambda_j + K'] \phi_j = 0$$  \hspace{1cm} (209)$$
assuming the independence of $\phi_j$, $j = 1, \cdots, 6n$ (previously proven). The eigenvalues from Eq. (209) are imaginary, and exist in complex conjugate pairs, i.e., $\lambda_j, \text{conjugate} = \pm i\sigma_j$.

Consider now the $6n \times 6n$ matrix

$$\phi = [\phi^1, \phi^2, \cdots, \phi^{6n}]$$  \hspace{1cm} (210)$$
and the transformation

$$q = \phi \eta$$  \hspace{1cm} (211)$$
The orthogonality relationships

$$\phi^k M' \phi^j = 0, \quad k \neq j$$  \hspace{1cm} (212)$$
$$\phi^k K' \phi^j = 0, \quad k \neq j$$

are well known. With suitable normalization of the eigenvectors, the condition

$$\phi^k M' \phi^k = 1, \quad k = 1, \cdots, 6n$$  \hspace{1cm} (213)$$
can be imposed. These relationships permit the transformation of Eq. (211) into Eq. (206) to provide (after premultiplication by $\phi^T$)

$$\ddot{\eta} + \sigma^2 \eta = \phi^T L'$$  \hspace{1cm} (214)$$
where, as previously,

$$\sigma^2 = \begin{bmatrix} \sigma_1^2 & 0 \\ \vdots & \ddots \\ 0 & \sigma_{6n}^2 \end{bmatrix}$$

Truncation of the modal coordinates $\eta_1, \cdots, \eta_{6n}$ to the set $\eta_1, \cdots, \eta_N$ is accomplished as previously, and symbolized by an overbar. Modal damping may also be incorporated, to obtain

$$\ddot{\bar{\eta}} + 2\xi \sigma \ddot{\bar{\eta}} + \sigma^2 \bar{\eta} = \phi^T L'$$  \hspace{1cm} (215)$$
or, more specifically (from 207),

$$\ddot{\bar{\eta}} + 2\xi \sigma \ddot{\bar{\eta}} + \sigma^2 \bar{\eta} =$$

$$-\ddot{\phi}^T M (\xi_{oo} - \xi_{eo} \ddot{H} - 7\xi_{eo}) \ddot{\eta} - \ddot{\phi}^T M E \bar{\eta}$$

$$+ \ddot{\phi}^T M \xi_{eo} m \ddot{E} \bar{\eta} + \ddot{\phi}^T \lambda$$  \hspace{1cm} (216)$$

The rationale for the incorporation of modal damping has been essentially physical; it has simply been observed from test data that dynamic simulation of a structure can be accomplished more accurately by inserting the matrix $\xi$ in Eq. (215) than by omitting it or guessing at an appropriate velocity coefficient matrix $D'$ to add $D' \dot{q}$ to Eq. (206). It is now possible to establish the mathematical significance of the assumption of modal damping. Imagine that Eq. (206) is modified by the incorporation of viscous-damping terms,

$$M' \ddot{q} + D' \dot{q} + K' q = L'$$  \hspace{1cm} (217)$$
and that $D'$ is a linear combination of $M'$ and $K'$,

$$D' = \alpha M' + \beta K'$$  \hspace{1cm} (218)$$
with $\alpha$ and $\beta$ scalars. The transformation $q = \phi \eta$ considered previously, with a premultiplication by $\phi^T$, then yields

$$\ddot{\eta} + (\alpha E + \beta \sigma^2) \ddot{\eta} + \sigma^2 \eta = \phi^T L'$$  \hspace{1cm} (219)$$
Since $\alpha E + \beta \sigma^2$ is a diagonal matrix, it does correspond to modal damping, with

$$\alpha + \beta \sigma^2 = 2\xi_i \sigma_j$$  \hspace{1cm} (220)$$
The two scalars $\alpha$ and $\beta$ may be chosen arbitrarily, and with the choice of these two numbers all values of the percentages of critical damping $\xi_1, \cdots, \xi_{6n}$ are established. (In engineering practice, more than two values of modal damping $\xi_1, \cdots, \xi_N$ are often prescribed independently, which involves a minor mathematical contradiction.)

Equation (216) is the final form of the appendage equation to be used in a space vehicle simulation when the appendage base is not supposed to rotate. Thus this is another alternative to Eqs. (168), (173), and (201). Equation (216) is to be used in conjunction with equations of motion of the vehicle, the rotor, and the damper, as well as the kinematic equations and control system equations required for a complete simulation. Because the base rotations are small, many of these equations simplify substantially. The vehicle rotational equation has...
already been recorded for this special case as Eq. (130). With the transformation \( q = \phi h \), this equation becomes

\[
T = 1* \ddot{h} + \dot{h} \ddot{h} + mb \ddot{E}^2 \\
+ (\Sigma_{\phi h} + \vec{R}E_{E0} + \Sigma_{\phi h} \vec{r}) M \dot{\phi h}
\]  

(221)
The damper equation (134) simplifies to

\[
m(1 - m/G_1) \ddot{x} + d \ddot{x} + k \dot{x} = \\
mE_{g1} [-F/G_1 + b \dot{E}^2 \ddot{E} + \Sigma_{\phi h} \mu \dot{\phi h}]
\]  

(222)
The rotor equation (136) becomes

\[
\tau = \beta (\dot{x} + E_{g1} \dot{E})
\]  

(223)

If the nominal value of the rotor angular momentum \( h \) relative to the base \( B_1 \) is \( h = \beta \dot{E} \), and the difference \( \Omega - \dot{\chi} \) is assumed to be small and is included in the linearization, the vehicle-rotation equation (221) becomes

\[
T = 1* \ddot{h} - \Omega \dot{E} \ddot{E} + \dot{\phi h} \ddot{E} + mb \ddot{E}^2 \\
+ (\Sigma_{\phi h} + \vec{R}E_{E0} + \Sigma_{\phi h} \vec{r}) M \dot{\phi h}
\]  

(224)
Equations (216), (222), (223), (224) are a complete system of dynamic equations for a dual-spin vehicle with a de-spun platform to which symmetric rotor, damper, and flexible appendage are attached. Vehicle simulation requires only the incorporation of control laws for \( \tau \) and \( T \).

7. Modal analysis of non-rotating structures using cantilever modes. As noted at the beginning of Section III-D, more than one useful coordinate transformation may be available for a given set of equations. The preceding equations of motion for a dual-spin vehicle with a de-spun platform provide an example. The final set of equations (216) and (222)-(224) were obtained by imposing the appendage coordinate transformation of Eq. (211) on a system of equations (207), (130), (134), and (136) that served as equations of motion of the appendage, the damper, the rotor, and the total vehicle in rotation. Note that the total vehicle translational equation (131) is not included explicitly in this system. The quantity \( \dot{\phi h} \) that characterized the vehicle-mass-center acceleration did appear originally in the equations of motion of the appendage (see Eq. 93), but this quantity was removed when the appendage equation was written in the form of Eq. (139), by substituting \( \dot{\phi} h = F/G_1 \) from Eq. (138). This step was motivated by the desire to separate the vehicle-trajectory problem from the attitude-dynamics problem, which is of paramount interest in this report. One could of course equally well retain the expression \( \dot{\phi} h \) in Eq. (139), and retain the translational equation \( F = \partial h \) in the simulation. This alternative does not preclude the selection of the transformation \( q = \phi h \) of Eq. (211), but it raises the possibility of another choice.

It should be noted that the decision to write the vehicle translational equation in terms of the coordinates \( X \) of the vehicle mass center \( CM \) was an arbitrary one. One could as well work with the inertial position vector (say, \( x \)) of the point \( O \), fixed in body \( B_1 \) and corresponding to the CM when the vehicle is in a nominal (undeformed) state. Then, in terms of previous notation (see Fig. 4),

\[
x = X + c
\]  

(225)
and the translational equation of the vehicle (Eq. 131) becomes

\[
F = [\dot{x} - \ddot{c} - \ddot{c} - 2 \ddot{c} - \ddot{c} - \ddot{c}] \partial h
\]  

(226)
where

\[
x = (i)^f x
\]  

(227)
In the linear approximation, with \( \omega = \dot{\theta} \), this is

\[
F = \partial h [\dot{x} - \ddot{c}]
\]  

(228)
The appropriate form for \( c \) can be obtained by linearizing Eq. (133) as

\[
\ddot{c} = -x_{E0} (M/\partial h) \ddot{E} - m \ddot{E}^2 \partial h
\]  

(229)
so the vehicle translational equation becomes

\[
F/\partial h = \dot{x} - x_{E0} (M/\partial h) \ddot{E} + m \ddot{E}^2 \partial h
\]  

(230)
Although it may seem unwise, it would not be incorrect to include Eq. (230) in the system simulation (just as \( F/\partial h = \dot{x} \) is carried along with the other equations of motion, except for the differences of coupling and complexity). Now if \( F/\partial h \) from Eq. (230) is substituted into the appendage equation (207), certain cancellations occur to provide

\[
M \ddot{E} + Kq = -M (x_{ax} - x_{ax} \vec{R}) \ddot{E} - M \ddot{E} \dot{q} + \lambda
\]  

(231)
This equation also has the form of Eq. (206), so the appendage coordinate transformation procedure applied to Eq. (207) applies as well to Eq. (231). The eigenvalues
and eigenvectors for these two sets of equations differ, of course, so the transformation in Eq. (211) is written as

\[ q = \phi^e \eta^e \]  

(232)

to provide a distinction. The transformed version of Eq. (231), after truncation and the introduction of damping, is

\[ \ddot{\eta}^e + 2\tau^e \dot{\eta}^e + (\sigma^e)^2 \eta^e = -\ddot{\phi}^e M (\Sigma_{cE} - \Sigma_{cE} \hat{R} - \hat{R} \Sigma_{cE}) \dot{\phi}^e - \ddot{\phi}^e M \Sigma_{cE} \theta^e + \ddot{\phi}^e \lambda \]  

(233)

In addition to this appendage equation, one must carry along equations of motion of rotor, damper, and vehicle rotation, which are the same as Eqs. (222)–(224) except that \( \ddot{\phi} \) becomes \( \ddot{\phi}^e \) and \( \theta \) becomes \( \eta^e \). One must also retain the vehicle translational equation (230), which appears when transformed as

\[ F/\eta = \delta \dot{\xi} + \Sigma_{cE}^T (M/\eta) \ddot{\phi}^e \eta^e + m \ddot{E}^t/\eta \]  

(234)

The change from modal coordinates \( \eta \) to modal coordinates \( \eta^e \) may appear to be of dubious merit, and indeed it does have disadvantages. It has apparently necessitated the retention of equations of translation in the attitude-control simulation of the vehicle, thus adding to the dimension of the system of equations being solved. Furthermore, this change violates the objective of seeking the transformation that uncouples the greatest number of equations, since the equations of translation newly added to the simulation (Eq. 234) are coupled in the modal coordinates \( \eta^e \).

The primary advantage of the alternative coordinates \( \eta^e \) (as opposed to \( \eta \)) stems from the convenience of their physical interpretation and experimental corroboration, and from the fact that any structural dynamics organization has the immediate capability of computing the eigenvalues and eigenvectors \( \lambda_m^e \) and \( \phi_m^e \), \( m = 1, \cdots, N \). To these pragmatic arguments may be added the fact that for most space vehicles the transformation \( q = \phi^e \eta^e \) is almost indistinguishable from \( q = \phi \eta \), and the added equation of motion (234) often does not actually have to remain in the simulation, because the terms involving \( \ddot{\eta}^e \) are small when the appendage is a small part of the vehicle mass. Finally, it can be argued (as will be shown) that when more than one appendage is attached to a given base, the coordinates \( \eta^e \) might simulate the appendage response for certain vehicle motions even better than the coordinates \( \eta \).

To properly evaluate the advantages and disadvantages of the alternative coordinate systems, one must make a physical interpretation. Both \( \eta_m \) and \( \eta_m^e \) are classical modal coordinates, in the sense that both measure the participation of the entire appendage structure in a vibration at a given frequency \( (\sigma_m \) or \( \sigma_m^e \) ), with all portions of the structure oscillating in phase as the structure deforms harmonically into a given mode shape \( (\phi_m \) or \( \phi_m^e \) ). The mode shapes and modal frequencies in question differ only in the boundary conditions imposed on the appendage.

The homogeneous counterpart to Eq. (231),

\[ M \ddot{q} + K q = 0 \]  

(235)

corresponds to the free vibrations of the appendage while on a fixed base. Accordingly, the modal coordinates in \( \eta \) are called the cantilever mode coordinates. Their significance is easily visualized, and the calculated mode shapes, frequencies, and damping ratios are easily confirmed by test.

The homogeneous form of Eq. (207) is

\[ M (E - \Sigma_{cE} \Sigma_{cE}^T M/\eta) \ddot{q} + K q = 0 \]  

(236)

This equation describes the free vibrations of the appendage attached to a base that is constrained against rotation, but free to translate as the base mass is pushed around by the shear forces at the base of the appendage. Note that the phrase “base mass” actually includes the mass of all of the vehicle except for the appendage in question (even if other appendages are present). Experimental corroboration of the mode shapes, frequencies, and damping ratios would be difficult to accomplish directly. In fact, when these coordinates are used, the modal data would probably not be obtained directly from Eq. (236). They would instead be computed by first applying to Eq. (236) the transformation \( q = \phi^e \eta^e \) to obtain

\[ \ddot{\eta}^e - \phi^e (M \Sigma_{cE} \Sigma_{cE}^T M/\eta) \phi^e \eta^e + (\sigma^e)^2 \eta^e = 0 \]  

(237)

and then after some truncation seeking a new transformation to diagonalize these equations. The cantilever modes would then be available from the preliminary transformation, and experimental corroboration would concentrate on verifying the cantilever mode data.

For vehicles with more than one appendage, it may happen that the cantilever modes simulate the vehicle behavior better than the modes with the base rotationally fixed and translationally movable. Consider for example the rigid body with two identical and symmetrically dis-
posed appendages shown in Fig. 10. The sketches in Figs. 10a, b, and c show the vehicle as it might actually respond if it were rotationally and translationally free. Among the many modes of vibration of which the vehicle is capable, the two shown in Figs. 10b and c involve symmetrical and antisymmetrical appendage deformations, respectively. In either vibration, the vehicle mass center must stay on the inertially fixed line shown, but for the symmetrical deformation in Fig. 10b the base-fixed point O moves, say, a distance ε. If now the appendage response is to be simulated by means of a single modal coordinate, either \( \eta^e \) or \( \eta \), one must judge the acceptability of these coordinates as the corresponding mode shape conforms to the actual deformations shown in Figs. 10b and c. This correspondence will depend on the faithfulness with which the assumed boundary conditions of the appendage match those of the actual vibrations. As illustrated in Figs. 10e and f, the cantilever mode permits no base translation, while the mode with base movable only in translation does allow the base to move an amount \( \delta < \varepsilon/2 \). Thus the boundary conditions shown in Fig. 10f provide mode shapes better suited than the cantilever modes to simulate the symmetrical vibration illustrated in Fig. 10b. Conversely, however, the cantilever modes, which permit no base motion, more accurately match the boundary conditions of the asymmetrical vibrations shown in Fig. 10c than do the modes in Fig. 10f. Thus, for a given truncation, the cantilever modal coordinates \( \eta^e \) are conceivably superior to the less constrained modal coordinates \( \eta \). This condition can occur only for multiple appendages, however. Since it is always possible to treat any number of structures attached to a given base as a single flexible appendage, the theoretical advantage seems always to lie with the modal coordinates corresponding to a rotationally fixed but translationally movable base. (The alternative method, which permits rotation as well as translation of the base, is pursued in Section IV.)

A practical compromise can be introduced that permits the pragmatically attractive use of cantilever modal coordinates without the addition of the vehicle translational equation into the simulation. There is no obstacle to the use of the transformation \( q = \phi^e \eta^e \) in Eq. (207). After multiplication by \( \phi^e \), one obtains Eq. (237), but with right-hand side \( \phi^e L' \). Although these equations are not uncoupled, because of the terms in \( -\phi^e (M \Sigma E_0 \Sigma \dot{E}_0^e M / \Sigma \dot{M}) \phi^e \eta^e \), these terms are generally small. Truncation can then be imposed without overwhelming concern in most cases, and the result is the appendage equation (with damping added)

\[
[E - \phi^e M \Sigma E_0 \Sigma \dot{E}_0^e (M / \Sigma \dot{M}) \phi^e] \ddot{\eta}^e + 2\phi^e \sigma^e \dot{\eta}^e + (\sigma^e)^2 \eta^e = \\
-\phi^e M (\Sigma E_0 - \Sigma \dot{E}_0 - \Sigma \dot{E}_0) \ddot{\theta} - \phi^e M \Sigma E_0 (F / \dot{M} - m\ddot{E}^e / \dot{M}) + \phi^e \lambda \tag{238}
\]

*This example was suggested by R. M. Bamford.
This appendage equation could be obtained equally well by replacing the expression $\omega x$ in Eq. (233) with its equivalent from Eq. (234).

E. Method of Synthetic Modes

At the beginning of Section III-C it is noted that there is a basic choice to be made in selecting the dynamic system for which equations of motion should be written in support of the flexible-appendage deformation equations developed in Section III-B. In the vehicle equations derived in Section III-C, the dynamic system chosen is the entire vehicle, including the flexible appendages. The vehicle equations, in conjunction with the appendage equations derived earlier, provide a basis for a complete dynamic simulation of the vehicle. The coordinate transformations presented in Section III-D are for the purpose of making the simulation equations more useful for practical computations.

Although the approach adopted in deriving the vehicle equations is always valid, the resulting equations may not in every case be in a form suitable for efficient computation. For this reason, equations are derived in this section that serve as alternatives to the vehicle equations of Section III-C. Whereas the equations of motion previously derived were for the entire vehicle, here the appendages are excluded from the system of rigid bodies for which equations of motion are written. The influence of the appendages on the motion of the rigid bodies is reflected as a contribution to the external resultant force and torque.

Equations of motion are to be derived here for a vehicle slightly less general than that illustrated in Fig. 9. The nutation damper $B_2$ is omitted here, since it would contribute terms to the equations that differ little from those derived in Section III-C. Only one flexible appendage is admitted, since the extension to two or more appendages is a trivial matter of repeating certain groups of terms. Relative rotation $\Omega$ of the appendage and its base is accommodated, since it is for vehicles in this class that the method of synthetic modes may prove advantageous.

Again the Newton-Euler equations of translation and rotation are to be derived for the dynamic system, only now this system consists merely of a rigid body $B_1$ in which a symmetric rotor $B_2$ has a fixed position and orientation. If $P$ is the inertial position vector of the mass center $P'$ of this system, $\Omega \eta'$ is the system mass, and the applied resultant force is the sum $f + f'$, with the latter accommodating all forces applied to $B_1$ by the appendage $A$, the translational vector equation is simply

$$f + f' = \Omega \eta' \dot{P}$$

In terms of the vector arrays $(b)$ and $(i)$, fixed respectively in $B_1$ and inertial space, and related by Eq. (53), this vector equation becomes

$$(b)\tau f + (b)\tau f' = \Omega \eta' (i)\tau \dot{P} = \Omega \eta' (b)\tau \dot{P}$$

which provides the matrix equation

$$\Omega \eta' \tau \dot{P} = f + f' \quad (239)$$

The rotational equation of the system $B_1$, plus $B_2$, may similarly be written as

$$I + I' = \frac{d}{dt} (I' \cdot \omega + h) \quad (240)$$

where the external torque about the system mass center $P'$ is separated into $I$ and $I'$, with the latter accommodating all torques applied to $B_1$ by the appendage $A$. Here $I'$ is the inertia dyadic for point $P'$ of the vehicle excluding the appendage, and $\omega$ and $h$ are as defined previously. Thus the expression $I' \cdot \omega + h$ is the sum of the angular momentum the system would have with a nonspinning rotor plus the relative angular momentum contribution of the rotor relative angular velocity. The inertia dyadic $I'$ is constant in basis $(b)$, so the differentiation of Eq. (240) yields only

$$I + I' = I' \cdot \dot{\omega} + \omega \times I' \cdot \omega + \dot{h} + \omega \times h \quad (241)$$

In matrix terms in basis $(b)$, with the definitions

$$I = (b)\tau I, \quad I' = (b)\tau I', \quad I' = (b)\tau I' (b) \quad (242)$$

Eq. (241) is equivalent to

$$I' \omega + \omega I' \omega + \dot{h} + \omega \times h = I + I' \quad (243)$$

The apparent simplicity of Eqs. (239) and (243), in comparison for example with the final equations of Section III-C (Eqs. 129 and 131), disappears when the appendage force $f'$ and torque $I'$ are calculated explicitly. These matrices must be proportional to the deformation of the appendage, which is assumed to be linearly elastic. The explicit expressions for $I'$ and $f'$ as linear functions of the deformation matrix $q$ are obtained somewhat circuitously in what follows.
Consider initially the forces and torques transmitted to the base of an elastic body A vibrating on an inertially fixed base. In this special case, the only inertial accelerations of the sub-bodies of the appendage are those due to vibrations, and the forces transmitted to the base are simply the inertial forces induced by vibration, i.e.,

\[ f' = (b)^T f' = - (a)^T \sum_{s=1}^{n} m^s \ddot{u}^s \]  

(244)

where (as previously) the vector basis \( (a) \) is fixed in the appendage base. Equation (50) provides \( (a)^T = (b)^T C^r \), so the matrix equivalent of Eq. (244) is

\[ f' = - C^r \sum_{s=1}^{n} m^s \ddot{u}^s \]  

(245)

The vector torque about point \( P' \) is then given by

\[ l' = (b)^T l' = - (a)^T \sum_{s=1}^{n} (m^s \dddot{r}^s + \ddot{p}^s) + (b)^T \ddot{p} \]  

(246)

where \( p = (b)^T p \) is the vector from point \( P' \) to point \( Q \) fixed on the interface between the appendage \( A \) and the rigid body \( B \). The matrix counterpart to Eq. (246) is

\[ l' = - C^r \sum_{s=1}^{n} (m^s \dddot{r}^s + \ddot{p}^s) - \ddot{p} C^r \sum_{s=1}^{n} m^s \ddot{u}^s \]  

(247)

In terms of the matrix \( q \) and the operators \( \Sigma_{\text{ex}} \) and \( \Sigma_{\text{in}} \) (see Eqs. 83 and 85), the expressions \( f' \) and \( l' \) can be written as

\[ f' = - C^r \Sigma_{\text{ex}} M \ddot{q} \]

\[ l' = - C^r (\Sigma_{\text{ex}} \dddot{r} + \Sigma_{\text{in}} M \ddot{q}) - \ddot{p} C^r \Sigma_{\text{ex}} M \ddot{q} \]  

(248)

These results apply only to the case of an appendage vibrating on a fixed base, and more general expressions are required for the problems of primary interest in this report. The necessary generalization is most easily accomplished after transformation of the appendage deformation coordinates \( q \) into modal coordinates. In the present restricted context of appendage vibration on a fixed base, the appendage vibration equations are the "cantilever" equations of Eq. (235), and the appropriate modal coordinate transformation is

\[ q = \phi^c \eta^c \]  

(249)

as in Eq. (232). The substitution of \( \ddot{q} = \phi^c \dddot{\eta}^c \) into Eqs. (248) provides an alternative expression for \( f' \) and \( l' \) for the special case of vibrations on a fixed base. This same substitution into Eq. (235) yields a solution for which

\[ \dddot{\eta}^c = -(\sigma^c)^2 \eta^c \]  

(250)

so that Eq. (248) may also be expressed as

\[ f' = C^r \Sigma_{\text{ex}} M \phi^c (\sigma^c)^2 \eta^c \]  

(251)

and

\[ l' = [C^r (\Sigma_{\text{ex}} \dddot{r} + \Sigma_{\text{in}}) + \ddot{p} C^r \Sigma_{\text{ex}}] M \phi^c (\sigma^c)^2 \eta^c \]  

(252)

Although Eqs. (248) are restricted to the fixed-base special case, and Eqs. (251) and (252) were obtained from Eqs. (248), still it can be argued that the expressions for \( f' \) and \( l' \) in Eqs. (251) and (252) are not so restricted, and indeed are completely general (for an elastic appendage). This follows from the necessary uniqueness of the force-displacement relationship of an elastic structure.

Thus Eqs. (251) and (252) may be substituted into Eqs. (239) and (243) to provide a general set of equations of motion of the rigid body to which the appendage is attached, as follows:

\[ M \dddot{q} + f + C^r \Sigma_{\text{ex}} M \phi^c (\sigma^c)^2 \eta^c \]  

(253)

\[ I' \dddot{\omega} + \dddot{\omega} + \dddot{\omega} = \]

\[ I + [C^r (\Sigma_{\text{ex}} \dddot{r} + \Sigma_{\text{in}}) + \ddot{p} C^r \Sigma_{\text{ex}}] M \phi^c (\sigma^c)^2 \eta^c \]  

(254)

These equations must of course be augmented by a rotor equation (e.g., Eq. 136), and perhaps also by control equations and kinematic equations. In addition, some form of the appendage equation is required. The final results of Section III-B would suffice (e.g., Eq. 84), or the individual sub-body equations of motion recorded as Eqs. (79) and (80) may be employed. Whichever are selected, the appendage equations must be subjected to the transformation \( q = \phi^c \eta^c \) for coordinate consistency.

The final system of equations of motion is useful in space vehicle simulation only if substantial truncation of the appendage deformation matrix \( \eta^c \) can be accomplished. This step cannot be undertaken casually in application to the equations of motion (253) and (254) of the rigid body to which the appendage is attached. The truncation rationale advanced in Section III-D favors the retention of the modal coordinates with the lowest frequency, and this policy often produces seriously deficient representations of the force \( f' \) and torque \( l' \) applied...
to the appendage base (except when the base is stationary). This fact is easily illustrated with a simple example.

Consider the vehicle of Fig. 11, which consists of a rigid body $B$ with a cantilevered elastic mast with a tip mass $m$ that greatly exceeds the mast weight. It is evident by inspection that the lowest-frequency cantilever-mode response involves transverse bending of the mast (see dashed lines of Fig. 11). A modal response of much higher frequency corresponds to vibration along the longitudinal axis of the mast. If this high-frequency mode is excluded in the modal-coordinate truncation process, the mast vibration response and the transmission of forces and torques to $B$ is faithfully represented if the body $B$ is inertially stationary. If, however, body $B$ is inertially accelerating, the force $f'$ and torque $l'$ are grossly misrepresented by this truncation (although an adequate representation of mast deformation may be preserved). Imagine for example that $B$ is accelerating at a constant rate $G$, along the centerline shown in the figure, so that the appendage reaches a steady-state small deformation after vibration attenuation. It is then obvious from first principles that the force $f'$ must be of magnitude $mG$ and must be directed along the indicated centerline of $B$, and the torque $l'$ must be of magnitude $mGe$ and direction normal to the plane of the paper, where $e$ is the eccentricity of $m$ as shown in the figure. Yet if truncation preserves only the transverse mode sketched in Fig. 11, the force $f'$ in Eq. (251) must be directed transversely to the mast, and the torque must be of magnitude $f'L$, where $L$ is the mast length. One might say that the truncated simulation is deficient only in its exclusion of the rigid-body modes of the mast. Although this may serve as a conceptual explanation of the problem, the difficulty is not rectified simply by adding rigid-body modal coordinates to the modal deformation coordinates in $\eta^e$, since this would unduly increase the interaction forces. One may instead devise “synthetic” modes for inclusion in $\eta^e$, as first suggested in Ref. 28. This procedure is briefly outlined here for a special case that serves to illustrate the method.

Consider a restricted problem of the dynamic simulation of a vehicle consisting of a rigid base $B$ and a flexible appendage $A$. Let the base $B$ be restricted to small deviations from a state of inertial rest, while the attached appendage $A$ rotates relative to $B$ at a small scanning rate.

The base equations (253) and (254) then become, in the linearized approximation,

$$\begin{align*}
\mathbf{\eta}' \ddot{\mathbf{P}} &= \mathbf{f} + C^T \Sigma_{E_0} M \Phi^e (\sigma^e)^2 \eta^e \\
\mathbf{I}' \ddot{\mathbf{\theta}} &= \mathbf{l} + [C^T (\Sigma_{E_0}^T \mathbf{f} + \Sigma_{O_0}^T) + \mathbf{p} C^T \Sigma_{E_0}^T] M \Phi^e (\sigma^e)^2 \eta^e
\end{align*}$$

(255)

(256)

In combination as a single matrix equation in the 6 by 1 matrix variable

$$\mathbf{U} = \begin{bmatrix} \mathbf{P} \\ \mathbf{\theta} \end{bmatrix}$$

(257)

these equations may be written as

$$\begin{align*}
\mathbf{J} \ddot{\mathbf{U}} &= \Lambda + \Sigma_{E_0} C^T \Sigma_{E_0}^T M \Phi^e (\sigma^e)^2 \eta^e \\
&+ \Sigma_{O_0} [C^T (\Sigma_{E_0}^T \mathbf{f} + \Sigma_{O_0}^T) + \mathbf{p} C^T \Sigma_{E_0}^T] M \Phi^e (\sigma^e)^2 \eta^e
\end{align*}$$

(258)

where the new symbols are

$$\begin{align*}
\mathbf{J} &= \begin{bmatrix} \mathbf{\eta}' \mathbf{f}^e & 0 \\
0 & \mathbf{I}' \end{bmatrix} \\
\Lambda &= \begin{bmatrix} f \\ l \end{bmatrix}
\end{align*}$$

It should be emphasized at this point that the number of rows in the matrix operators denoted by $\Sigma_{E_0}$ and $\Sigma_{O_0}$ is
established solely by context (see Eq. 85). Because the matrix $C^T$ in Eq. (258) is 3 by 3 and the matrix $\Lambda$ is 6 by 1, the premultiplier $\Sigma_{EE}$ must be of dimension 6 by 3. The premultiplier $\Sigma_{EE}^T$ of the 6n by 6n matrix $M$ must, however, be of dimension 3 by 6n. It is conceivable that a symbol such as $\Sigma_{EE}$ may appear twice in one equation and represent two matrices of different dimension, as established by context.

In conjunction with Eq. (258) for body $B$, one must consider the appropriate appendage equations of motion. These must be constructed from the ingredients of Eqs. (79) and (80) for the appendage sub-bodies, following a pattern established in detail in Section III-B, and culminating in Eq. (84). Present interest is restricted to the case of small $\omega \equiv \theta, \epsilon \equiv 0$, and variable $C$. Appendage equations of the form of the cantilever equations of Eq. (231) are sought, with added terms as necessary to accommodate the slowly changing direction cosine matrix $C$ (or the corresponding angular velocity $\Omega$). In Eqs. (79) and (80), the only added terms that survive linearization are $I_1 \dot{\psi}^2$ and $-m \dot{\theta} \dot{\psi}^2$, respectively. Thus the necessary modification of appendage equation (231) yields (with $\bar{P}$ replacing $\delta x$ and $\bar{P}$ replacing $\bar{R}$)

$$M \ddot{q} + Kq = -M (\Sigma_{EE} C - \Sigma_{EO} C \bar{P} - \bar{\gamma}\Sigma_{EO}) \dot{\theta}$$
$$- M \Sigma_{EO} C \bar{P} + \lambda - M (\Sigma_{EE} - \bar{\gamma}\Sigma_{EO}) \dot{\theta}^2$$

(259)

A complete derivation of this equation may be found in Ref. 28, although notational differences must be reconciled to obtain confirmation. In terms of the matrix $U$ of Eq. (257) and the cantilever modal coordinates $\eta^c$ of Eq. (249), the appendage equation is (with modal damping included)

$$\ddot{\eta}^c + 2 \xi^c \dot{\eta}^c + \eta^c \ddot{\eta}^c = -\phi^c M (\Sigma_{ab} C - \Sigma_{EO} C \bar{P} - \bar{\gamma}\Sigma_{EO} C) \Sigma_{ab} \dot{U}$$
$$- \phi^c M \Sigma_{EO} C \Sigma_{EO} \dot{U} + \phi^c \lambda - \phi^c M (\Sigma_{EO} - \bar{\gamma}\Sigma_{EO}) \dot{\theta}^2$$

(260)

Equations (258) and (260) constitute a complete set for dynamic simulation, requiring only the specification of external environment or control law for the explicit determination of $\Lambda$, $\lambda$, $C$, and $\dot{\theta}^2$. Inspection of these equations reveals the repeated presence of a 6n by 6 matrix, here defined as $\Delta$ (somewhat different notation is employed in Refs. 28 and 29), as given by

$$\Delta = -\phi^c M (\Sigma_{EO} C \Sigma_{EO} + \Sigma_{EO} C \Sigma_{EO}^T - \bar{\gamma}\Sigma_{EO} C \bar{P} \Sigma_{EO}^T)$$

(261)

In terms of the matrix $\Delta$, the system equations of motion (258) and (260) become

$$J \ddot{U} = -\Delta^T (\sigma^e)^2 \eta^e + \lambda$$

(262)

and

$$\ddot{\eta}^e + 2 \xi^e \dot{\eta}^e + (\sigma^e)^2 \eta^e =$$
$$\Delta \dot{U} + \phi^c \lambda - \phi^c M (\Sigma_{EO} - \bar{\gamma}\Sigma_{EO}) \dot{\theta}^2$$

(263)

These equations are in a form well suited for practical simulation studies only after truncation of coordinate $\eta^c$ has been accomplished. As previously noted and illustrated with the example of Fig. 11, coordinate truncation must be undertaken very cautiously if valid representations of interaction forces and torques are to be preserved.

Coordinate truncation is greatly facilitated by the physical interpretation of the elements of the matrix $\Delta$. Corresponding to each modal coordinate in $\eta^c$ there is one row of six scalars in the matrix $\Delta$; i.e., the $j$th row of $\Delta$, consisting of $\Delta_{j1}, \ldots, \Delta_{j6}$, corresponds to the $j$th modal coordinate $\eta_j^c$. From Eq. (263), the steady-state modal response to a constant base acceleration $\dot{U}$ can be found to be

$$\eta^c = [(\sigma^e)^2]^{-1} \Delta \dot{U}$$

(264)

For the $j$th mode, the steady-state deformation is therefore

$$\eta_j^c = \sigma_j^e \sum_{i=1}^6 \Delta_{ji} \dot{U}_i$$
$$= \sigma_j^e \left[ \Delta_{j1} \dot{P}_1 + \Delta_{j2} \dot{P}_2 + \Delta_{j3} \dot{P}_3 + \Delta_{j4} \dot{\theta}_1 + \Delta_{j5} \dot{\theta}_2 + \Delta_{j6} \dot{\theta}_3 \right]$$

(265)

Substitution of Eq. (264) into (262) yields further physical interpretation. The result

$$J \ddot{U} = -\Delta^T \Delta \dot{U} + \lambda$$

(266)

provides in the 6 by 6 matrix $-\Delta^T \Delta$ a collection of the forces and torques (about $P$) applied by the appendage to the base due to a constant base acceleration. Consequently, the scalars $\Delta_{ji}$ have been called "dead-load coefficients." The 6 by 6 matrix $\Delta^T \Delta$ is evidently just a
collection of inertia-like quantities. As a matrix of 3 by 3 partitions, $\Delta^T \Delta$ may be written in the form

$$\Delta^T \Delta = \begin{bmatrix} \Omega^T \Omega & 0 \\ 0 & I^4 \end{bmatrix}$$

(267)

where $\Omega^T \Omega$ is the mass of the appendage, $I^4$ is the inertia matrix of the appendage with respect to point $P'$, and $p^T$ is the matrix in basis $\{a\}$ of the vector from $P'$ to the appendage mass center. Note that the matrix $\Delta^T \Delta$ is symmetric (since the transpose of the skew-symmetric matrix $\Omega^T \Omega$ is its negative).

With this simple physical interpretation of the matrix $\Delta^T \Delta$ comes a new rationale for coordinate truncation. If a valid representation of interactions is to be preserved, it is essential that the truncated $N$ by 6 matrix $\tilde{\Omega}$ obtained by replacing $\phi^c$ by $\tilde{\phi}^c$ in Eq. (261) continue to satisfy Eq. (267), i.e., that even after truncation $\tilde{\Delta}^T \tilde{\Delta} \approx \Delta^T \Delta$. In practical applications, this may be a difficult stipulation to meet; there may simply not be available among the eigenvectors (modal columns) in $\phi^c$ a small number of mode shapes that meet simultaneously the constraint of Eq. (267) and the earlier requirements that low-frequency modes and potentially resonant modes be retained. A pragmatic reaction to this dilemma is the creation of a valid representation of interactions is to be preserved, a small number of synthetic modes to the truncated coordinate matrix as necessary to obtain satisfactory correlation with the known right side of Eq. (267), without even confronting the necessity of solving simultaneous algebraic equations. If the six dead-load coefficients of the first synthetic mode are chosen so as to provide exactly the correct first column for $\tilde{\Delta}^T \tilde{\Delta}$, then the first of the six dead-load coefficients of the second mode must be zero, and the remaining five can be chosen to match the second columns of $\tilde{\Delta}^T \tilde{\Delta}$ and $\Delta^T \Delta$. This procedure continues, each time with one or more zero dead-load coefficients, until six synthetic modes have been added and independent elements of $\tilde{\Delta}^T \tilde{\Delta}$ and $\Delta^T \Delta$ match perfectly. Alternatively, one may mechanically record the 21 independent equations available from Eqs. (267) and (268), and solve simultaneously for 21 unknown dead-load coefficients. This would require only 4 synthetic modes (24 dead-load coefficients, with 3 arbitrarily assigned).

The final equations of motion for vehicle simulation then become

$$J\ddot{\mathbf{U}} = -\Delta^T (\ddot{\sigma})^s \ddot{\eta}^c + \Lambda$$

(269)

and

$$\ddot{\eta}^c + 2\bar{\varphi}^c \sigma^c \dot{\eta}^c + (\sigma^c)^2 \eta^c = \Delta \ddot{\mathbf{U}} + \bar{\varphi}^c \lambda - \bar{\varphi}^c \mathbf{M} (\mathbf{z}_{\text{geo}} - \mathbf{r}_{\text{geo}}) \dot{\eta}^a$$

(270)

where the barred matrices represent truncations that may include synthetic modes. It may be computationally advantageous to use Eq. (269) to rewrite Eq. (270) in the form

$$\ddot{\eta}^c + 2\bar{\varphi}^c \sigma^c \dot{\eta}^c + [(\sigma^c)^2 + \bar{\varphi}^{-1} \Delta^T (\ddot{\sigma}^c)] \eta^c = \Delta \ddot{\mathbf{U}} + \bar{\varphi}^c \lambda - \bar{\varphi}^c \mathbf{M} (\mathbf{z}_{\text{geo}} - \mathbf{r}_{\text{geo}}) \dot{\eta}^a$$

(271)

If the appendage response is of primary interest, Eq. (271) will suffice, but for space vehicle simulation, both Eqs. (269) and (271) are required.

The most significant feature of these equations is the simplicity of the coefficient matrices of the second derivative terms. By multiplying Eq. (269) by the inverse of the constant and usually diagonal matrix $J$, one can obtain Eqs. (269) and (271) in the combined matrix form
The absence of a coefficient matrix for the highest-ordered derivative simplifies the numerical integration of Eq. (272) very substantially when compared to corresponding equations of motion obtained by the methods of Section III-C. The assumption $C = E$ was introduced in the course of the derivation, and the final system of equations is not directly applicable to vehicles that undergo large changes of configuration. Had this feature (variable $C$) of the present equations been retained, the vehicle equations, such as Eq. (224), would have adopted a form that would be very inefficient for numerical integration. The matrix $1^*/E$ in Eq. (224) is the total vehicle inertia matrix, which would become a function of time, and the variable direction cosine matrix $C$ would remain in the coefficient matrix of $\dot{U}$. As a result, it would become necessary to invert or apply Gaussian elimination to a time-varying coefficient matrix of the highest-ordered derivative at each step of the numerical integration (or at greater intervals as judgment allowed). It may therefore be concluded that the method of this subsection (and of Ref. 28) is apt to be preferable in application to vehicles of configuration varying substantially with time.

\[
\begin{bmatrix}
\dot{U} \\
\dot{\theta}^e
\end{bmatrix} + \begin{bmatrix}
0 & 0 \\
0 & 2\dot{\theta}^e\alpha^e
\end{bmatrix} \begin{bmatrix}
\dot{\theta} \\
\dot{\theta}^e
\end{bmatrix} + \begin{bmatrix}
0 & \delta^e (\alpha^e)^2 \\
0 & \delta^e (\alpha^e)^2 + [E - \delta J^{-1} \delta^e] (\alpha^e)^2
\end{bmatrix} \begin{bmatrix}
U \\
\dot{\theta}^e
\end{bmatrix} = \begin{bmatrix}
\lambda \\
\delta J^{-1} \lambda + \delta^e \alpha^e M (\Sigma_{ae} - \Sigma_{be}) \dot{\theta}^e
\end{bmatrix}
\] (272)

The absence of a coefficient matrix for the highest-ordered derivative simplifies the numerical integration of Eq. (272) very substantially when compared to corresponding equations of motion obtained by the methods of Section III-C. The assumption $C = E$ was introduced in the course of the derivation, and the final system of equations is not directly applicable to vehicles that undergo large changes of configuration. Had this feature (variable $C$) of the present equations been retained, the vehicle equations, such as Eq. (224), would have adopted a form that would be very inefficient for numerical integration. The matrix $1^*/E$ in Eq. (224) is the total vehicle inertia matrix, which would become a function of time, and the variable direction cosine matrix $C$ would remain in the coefficient matrix of $\dot{U}$. As a result, it would become necessary to invert or apply Gaussian elimination to a time-varying coefficient matrix of the highest-ordered derivative at each step of the numerical integration (or at greater intervals as judgment allowed). It may therefore be concluded that the method of this subsection (and of Ref. 28) is apt to be preferable in application to vehicles of configuration varying substantially with time.

F. Summary

The discussion of hybrid-coordinate methods includes a very substantial body of material, much of which is not available elsewhere. In this respect, Section III differs from Sections II and IV, which are intended to provide abbreviated reviews of the well-established methods that employ either discrete coordinates or vehicle normal-mode coordinates exclusively.

To summarize, two quite different methods are developed: Section III-E covers the synthetic-mode method, which employs equations of motion written separately for the rigid and elastic components of the vehicle; and the preceding sections explore the various ways in which the equations of motion of the total vehicle may be combined with appendage equations in the simulation. Except in unusual circumstances, the synthetic-mode method should be applied only to vehicles with time-varying configuration, because this method generally requires more appendage modal coordinates than are necessary with the alternative approach.

Sections III-B and III-C contain derivations of equations of motion of flexible appendages and total vehicles, respectively. These derivations follow from the most basic principles of classical mechanics, but they are complex in detail. Vehicles of variable gross configuration are excluded in the course of the derivations of Section III-C, so that the most general final results of that section are applicable to the unrestricted motion of a vehicle consisting of a rigid body $B$, to which there are attached a rigid symmetric rotor, a linear oscillator, and a flexible appendage limited to small deformations. The appropriate equations are Eqs. (129), (131), (134), and (136), which may be considered in combination with Eq. (95) for the appendage deformations. Equation (84) provides the appendage equations of motion in a more general case in which the appendage is undergoing large rotations relative to its base.

The derivation of equations of motion for flexible vehicles in terms of discrete coordinates is principally a bookkeeping task. The equations of Section III-B and III-C become useful only after the coordinate transformations of Section III-D are imposed. Four distinct coordinate transformations are developed as shown in Eqs. (159), (187), (211), and (232). The first two are transformations to be applied to first-order equations, which must be used if modal coordinates (in the broadest sense) are to be used for an appendage that is either on a rotating base or subject to discrete damping. The various ways in which first-order transformations can be used, with discussion of limitations and advantages, may be found following Eq. (201). Second-order transformations (as in Eqs. 211 and 232) are evaluated in the last part of Section III-D.

Under certain conditions of engineering interest, it may be appropriate to apply the second-order equation modal transformations described here even to rotating appendages, although in general such application is restricted to nonrotating flexible bodies.
Motivated by the problem of dynamic analysis of helicopter blades, Bisplinghoff, Ashley, and Halfman provide in their well-known text (Ref. 13) a derivation of the equations of motion of a rotating beam (pp. 95–98) and the corresponding modal analysis (pp. 184–187). Their equations of motion (Eqs. 3-121 and 3-127 of Ref. 13) apply to a continuous beam rather than to the discretized structural model adopted in this report, but they differ from the equations of motion of a rotating appendage as derived in this report (Eqs. 143 and 140) in a more fundamental way also. The term \( G'q \) in Eqs. (143) and (140) does not have a counterpart in the equations of Ref. 13. The mathematical significance of this difference is reflected in the comparison of the modal analyses of Ref. 13 and this report. In Ref. 13 it remains possible to employ in modal analysis the class of second-order point transformations, which in this report are restricted to nonrotating systems (typified by Eqs. 211 and 232). According to Section III-D, one must employ first-order equation transformations (Eqs. 159 and 187) to inhomogeneous equations for rotating appendages, and may employ only contact transformations (Eq. 186) to second-order homogeneous equations of rotating appendages. These differences can be reconciled by physical interpretation of the significance of the term \( G'q \) in Eqs. (143) and (140).

For an undamped system, velocity-proportional terms must arise in application to a spinning structure due to Coriolis "forces" or Coriolis accelerations, the latter given by \( 2\omega \times v \), with \( \omega \) the angular velocity of the reference frame with respect to which \( v \) is the relative velocity. Because \( G'q \) comes from this vector cross-product, the matrix \( G' \) must be skew-symmetric. If, however, the structure is very stiff in directions pointing radially from the spin axis, so that \( v \) has no component in this direction, then the Coriolis forces are limited for small deformations to the radial direction. Since the structure is stiff in this direction, these forces may be of negligible influence on the deformation. It is quite a separate matter to ignore the influence of these forces on the reaction at the base of the structure, but this too is often a reasonable assumption.

In application to helicopter blades, it may be appropriate that Coriolis accelerations be ignored, as implied in Ref. 13. Etkin and Hughes (Ref. 38) have successfully applied the methods of Ref. 13 to a class of spin-stabilized satellites with radially directed tubular metal antennas, as typified by Alouette I and Explorer XX. Although reasonable caution must be exercised in application to very long, slender structures (since the Coriolis force applies a column load), the simplification of the coordinate transformation afforded by ignoring Coriolis forces should be recognized and this step should be considered carefully in any engineering analysis.

IV. Vehicle Normal-Mode Coordinate Methods

A. Application to Nongyroscopic Linear Systems With Structural Damping

In the introductory remarks of Section III-C, it is noted that the primary test of the utility of a given coordinate transformation is the degree to which it permits the truncation of the coordinate matrix. Thus the desirability of a given coordinate transformation should be measured in terms of the degree to which it uncouples the system of differential equations employed in the vehicle simulation. Yet in all of Section III, transformations are applied to the appendage deformation coordinates \( q \) only. This represents a compromise with the objective of uncoupling the equations, since in many applications coordinate transformations can be applied as well to the variables representing the attitude and position of the appendage base, so that even more complete uncoupling of equations results.

In this section, attention is directed to the determination of the range of applicability of point transformations of second-order equations in obtaining uncoupled or normal modal coordinates for the entire vehicle. Examples of transformations in this class are the appendage transformations of Eqs. (211) and (232).

In the discussion of Eq. (217), it is noted that any equation of the class

\[
M' \ddot{q} + D' \dot{q} + K' q = L'
\]

where \( M' \) and \( K' \) are symmetric and

\[
D' = \alpha M' + \beta K'
\]

with \( \alpha \) and \( \beta \) arbitrary scalars, permits the normal-mode transformation of Eq. (211), namely,

\[
q = \phi \eta
\]

where \( \phi \) is a square matrix whose columns are the eigenvectors associated with Eq. (373). With this transformation and a premultiplication by \( \phi^T \), Eq. (273) becomes

\[
\ddot{\eta} + 2\zeta \omega \dot{\eta} + \omega^2 \eta = \phi^T L'
\]

This represents a compromise with the objective of uncoupling the equations, since in many applications coordinate transformations can be applied as well to the variables representing the attitude and position of the appendage base, so that even more complete uncoupling of equations results.
where $\zeta$ and $\sigma$ are diagonal matrices containing, respectively, the percentages of critical damping and the natural frequencies of the modal coordinates in $\eta$.

This discussion applies as well to a matrix equation formed by combining the appendage equation with the remaining equations of motion of the vehicle. For example, one might consider Eqs. (207), (130), (131), (223), and a form of (134) obtained by linearizing in $\dot{\omega} \approx \dot{\omega}$. These equations constitute a complete system of dynamic equations for a dual-spin vehicle with a de-spun platform to which symmetric rotor, damper, and flexible appendage are attached. With the hybrid-coordinate approach, the transformation of Eq. (275) is successfully applied to the appendage deformations, and the resulting system of equations is recorded as Eqs. (216), (222), (223), and (224). Now, however, by virtue of the term $-\dot{h}\dot{\omega} = -\beta\Omega E\dot{\omega}$ in Eq. (130) and the term $d_2\xi$ in Eq. (134), the indicated system of equations of the entire vehicle does not have the structure of Eq. (273), and transformation (275) is no longer useful.

Only by removing the rotor and the damper from the vehicle can the system of equations be made to conform to Eq. (273). Then the total system of dynamic equations reduces to

$$M\left(E - \Sigma_{E\theta} \Sigma_{E\theta}^T M/\Omega n\right) \ddot{\eta} + K\eta = -M \left(\Sigma_{E\theta} - \Sigma_{E\theta} \ddot{R} - \ddot{\tau}_{E\theta}\right) \dot{\omega} - M \Sigma_{E\theta} F/\Omega n + \lambda$$

(277)

and

$$T = I^* \ddot{\omega} + \left(\Sigma_{E\theta} + \ddot{R} \Sigma_{E\theta}^T + \Sigma_{E\theta} \ddot{\tau}\right) M \dot{\eta}$$

(278)

These equations may be written as the single matrix equation

$$ \begin{pmatrix} I^* & \Sigma_{E\theta} + \ddot{R} \Sigma_{E\theta}^T + \Sigma_{E\theta} \ddot{\tau} \end{pmatrix} M \begin{pmatrix} \ddot{\eta} \\ 0 \\ 0 \end{pmatrix} + \begin{pmatrix} 0 & 0 \\ 0 & K \end{pmatrix} \begin{pmatrix} \dot{\eta} \\ q \end{pmatrix} = \begin{pmatrix} T \\ \lambda - M \Sigma_{E\theta} F/\Omega n \end{pmatrix}$$

(279)

which has the structure of Eq. (273), lacking the damping term $D'$. One could equally well replace $F/\Omega n$ by $\Theta X$ and add Eq. (131) to the system of equations, extending the matrix of unknowns to include $\Theta X$. Alternatively (and equivalently), Eqs. (230), (231), and (278) could be combined as a single matrix equation.

Determination of eigenvalues and eigenvectors for Eq. (279) is conceptually straightforward, although computational obstacles may be introduced by the zero eigenvalues, which are a consequence of the positive semi-definiteness of the coefficient matrix of $[\dot{\omega}, \eta]^T$. Aside from the question of computational procedure, which is treated in any modern structural dynamics text (e.g., Ref. 11), there is the question of interpreting the physical significance of these zero-frequency modes. These modes correspond to translation of the mass center and rotation of the undeformed vehicle. When response in these modes is combined with response in the various deformation modes, it becomes somewhat difficult to determine just what it is that rotates in the amount given by the zero-frequency rigid-body-rotation mode. Milne treats this question in Ref. 4, pointing out that one might reasonably be interested in any of three reference frames that participate in the gross motion of the vehicle: (1) an attached reference frame, (2) a reference frame in which the mass center and the principal axes of the deforming vehicle are fixed, and (3) a "mean motion" frame with respect to which the vehicle mass center is fixed and vehicle relative motions have no angular momentum about the vehicle mass center. Milne provides a simple example that illustrates the possible differences in frames (2) and (3), and demonstrates that the zero-frequency rigid-body-rotation mode provides the rotation of the mean motion frame (3). (This fact is recognized also in Ref. 3, although less explicitly.)

Equation (279) is not quite as general as Eq. (273), because of the absence of damping. In order that the vehicle rigid-body modes have zero damping as well as zero frequency, any damping included must be proportional to stiffness, i.e., $D' = \beta K'$ is required. Within this restriction, which is traditionally acceptable in structural dynamics, one may transform a version of Eq. (279) with structural damping into the form of Eq. (276). Coordinate truncation may then be imposed as argued previously.

The term non-gyroscopic is applied to the linear systems of Eqs. (279) and (273) to reflect the absence of the term $G'\dot{q}$ (skew-symmetric $G'$) induced by rotation of the vehicle or some vehicle component.

Point transformations to normal-mode coordinates for a system of second-order equations for an entire space
vehicle are thus limited in application to nongyroscopic linear systems with structural damping. Physical application is thus limited to an inertially stabilized space vehicle with nonrotating appendages and no internal rotors or discrete dampers.

It should be noted that even now the original objective of uncoupling the entire system of equations is not realized, since in general the dynamic equations in Eq. (279) must be augmented by an equation that specifies any control torques and forces in $T$, $\lambda$, and $F$. The control equations are almost invariably nonlinear, if only due to a dead band in the response of thrusters to sensors, or the response of sensors to motion. The equations may also be "damped" (positively or negatively). Frequently the control equations are of higher order than Eq. (279). It is therefore unlikely that the control equations will be amenable to inclusion in a system of equations of the structure of Eq. (273), so they must remain as auxiliary equations that continue to couple the system of equations even after the transformation to vehicle normal-mode coordinates.

It may also be noted that even for passive systems the external torques $T$ and $\lambda$ may depend explicitly on the variables in $\theta$ and $q$. If, for example, gravity torque expressions are substituted for $T$ and $\lambda$, these terms should be shifted to the left side of Eq. (279) and their presence reflected in the modal analysis (with the resulting replacement of the zero-frequency modes by satellite libration modes). If this step is not taken, the terms $T$ and $\lambda$ (buried in $L'$) will introduce modal-coordinate coupling in Eq. (276).

B. Application to Gyroscopic Linear Systems With Discrete Damping

Equations of motion of a dual-spin vehicle with a de-spun platform with attached rotor, damper, and flexible appendage are collected as Eqs. (207), (130), (131), (223), and a linearized version of (134). These are all linear equations, but they are gyroscopic and involve discrete damping (so Eq. 274 is violated). Thus it is not possible to apply a point transformation to these second-order equations to obtain uncoupled modal coordinates.

There remains the possibility of rewriting the entire system of equations as a single linear state equation such as Eq. (157), i.e.,

$$\dot{Q} = BQ + L$$  \hspace{1cm} (280)

As noted in Section III-D in the development following Eq. (145), there is no obstacle to the application of a transformation as in Eq. (159), i.e.,

$$Q = \phi Y$$  \hspace{1cm} (281)

to obtain uncoupled equations in the modal coordinates in $Y$ (see Eq. 160). The disadvantages of this approach (stemming principally from complex numbers in $\phi$) are discussed extensively in Section III-D.

When the total system of second-order equations is gyroscopic, but no discrete damping is included, so that the homogeneous equations have the structure of Eq. (143), it may be preferable to employ the real transformation of Eq. (187)

$$Q = PZ$$  \hspace{1cm} (282)

to the state equation (280). An evaluation of the advantages of this transformation over that of Eq. (281) appears in the text following Eq. (204) in Section III-D.

The method of Foss (Ref. 36 and Eq. 173) may be applied to the entire system of vehicle equations in application to nongyroscopic but discretely damped systems.

There remains the question of the existence of normal-mode coordinates for an entire system of vehicle coordinates when the base is not inertially stabilized. When the base to which the appendage is attached has a substantial inertial angular velocity $\omega$, the relatively simple Eq. (130) for the vehicle rotation is replaced by its complicated antecedent, Eq. (129). This equation is nonlinear, so it will not yield directly to any kind of modal-coordinate transformation. In many problems of space vehicle control, however, one has advance knowledge of the desired behavior $\omega(t)$ of $\omega$. It is then possible to introduce the variational coordinates $\theta_1, \theta_2, \theta_3$, which define the small deviation of the vehicle from its nominal attitude, and to replace $\omega$ in Eq. (129) by

$$\omega = \omega + \dot{\theta}$$  \hspace{1cm} (283)

Assuming $\dot{\theta}$ to be small and including this with $q$ and $\xi$ in the linearization process, one can obtain from Eq. (129) a corresponding linear equation. When $\omega(t)$ depends on time, this equation will still not yield to modal-coordinate transformation, but when $\omega$ is constant, modal coordinates for the entire vehicle can be found. With careful bookkeeping, the resulting linear, constant-coefficient second-order differential equation obtained from Eq. (129)
can be combined with the correspondingly transformed and linearized Eqs. (134), (136) and (95) for damper, rotor, and appendage, and written as a state equation as in Eq. (280). Application of the transformation of Eq. (281) is then possible. All this would require much labor, and the end result would be of dubious value in comparison with the hybrid-coordinate methods of Section III.

C. Component Modal-Combination Methods

The practical implementation of some of the transformations discussed here and in Section III may for complex vehicles require modal analyses that strain the capacity of present digital computers. To circumvent the problem of finding eigenvalues and eigenvectors for the matrix equations of very high dimension that may be required to accomplish the simulation of an entire space vehicle, Hurty (Refs. 9, 10) has developed a method whereby separate modal analyses are performed on components or subsystems of the vehicle, and then a vehicle modal analysis is accomplished by combining component modes. Computer programs have been written for this purpose (e.g., Ref. 39), and the component modal-combination approach has been widely adopted.

It may appear from the discussion in this section that the use of vehicle modal coordinates is qualitatively different from the use of hybrid coordinates (Section III); the former method involves coordinate truncation for fully uncoupled dynamic equations, while the latter imposes truncation on appendage deformation coordinates in equations that are coupled by the discrete coordinates of the vehicle. As a practical matter, however, the use of vehicle modal coordinates often depends upon acceptance of the component modal-combination approach, and this method also involves the truncation of modal coordinates for components without regard for coupling of the component equations with those of the remainder of the vehicle. Thus the hybrid-coordinate approach and the component-mode method of modal analysis share the same pragmatic philosophy. In fact, it may develop in many applications that the hybrid-coordinate approach is a variant of the component-mode method in which certain components are assumed to be rigid and the final modal analysis of the entire vehicle is not undertaken.

D. Summary

This brief discussion of vehicle modal-coordinate methods is included for the primary purpose of keeping the hybrid-coordinate methods of the Section III in perspective. It has been observed that the same kinds of coordinate transformations that prove useful in application to appendage deformation coordinates can be applied to the entire system of vehicle coordinates.

Section IV-A contains a description of the dynamic systems to which the traditional second-order equation point transformations of structural dynamics may be applied. It has been noted that these simple modal-coordinate transformations are inapplicable when the vehicle contains any spinning parts or discrete devices that dissipate (or create) mechanical energy. These same transformations will still be applicable to the appendage coordinates, providing that the appendage base is not rotating.

In Section IV-B, modal-coordinate transformations are identified that are applicable to any linear dynamic system. These are transformations to be applied to first-order equations, and in some cases complex numbers are introduced by the transformation.

Although it is not the purpose of this report to explore methods of numerical computation, the component modal-combination method is briefly described in Section IV-C for comparison of its underlying coordinate-truncation philosophy with that of the hybrid-coordinate method.

V. Control System Simulation

A. Nonlinear System Analysis

Space vehicle attitude-control systems are very rarely linear in their relationship between control torque and attitude error. There is almost always a dead-band range within which the error can fall without actuating control torque devices, and usually nonlinearities are present even beyond the dead-band range. Often the torqueing devices are gas jets or other mechanisms operating in a pulsed or "bang-bang" mode, or they may be momentum storage devices that require periodic "momentum dumping." The sensors may be of the sampled data type, they may be inertial sensors that saturate beyond certain error limits, or they may be optical sensors with a limited linear range. Thus there are often significant nonlinearities in the sensors and actuators of a space vehicle attitude-control system, as well as the nonlinearities introduced by the logic of the control law.

The dynamic equations of the controlled vehicle are also frequently nonlinear, as the equations derived in the preceding sections attest, even when the structure under control is assumed to be linearly elastic.
Techniques of nonlinear analysis have been devised that permit practical application to scalar second-order differential equations, or to equations with certain kinds of nonlinearities. These methods are described extensively in Refs. 40 and 41, and they will not be reviewed here. It is quite possible that, with severely restrictive assumptions, equations for a nonlinearly controlled flexible space vehicle could be obtained for which nonlinear analysis may be fruitful.

For example, if for a given vehicle it is reasonable to assume a single-axis response to a given control torque, say, \( \theta_1 \neq 0 \) and \( \theta_2 = \theta_3 = 0 \), then a single second-order equation for \( \theta_1 \) may be extracted from the vehicle equations and subjected to phase-plane studies (Ref. 40, Chap. 7, and Ref. 41, Ch. 7). Alternatively, such simplified equations may for certain kinds of nonlinearities be amenable to quasi-linearization and the application of describing functions (Ref. 40, Chap. 9, and Ref. 41, Chap. 3).

Certainly the primary method of nonlinear analysis of a space vehicle with flexible appendages is direct simulation and integration of equations of motion. The modal analysis required for the explicit expression of the equations of motion of Sections III and IV is clearly suited to digital computer implementation, and in most cases this would appear to be true also of the integration process. With severe truncation of appendage modal coordinates, however, the equations may be efficiently employed in an analog simulation.

As indicated by the references cited in Section II, there is now abundant experience in the digital computer numerical integration of discrete coordinate equations of motion.

Hybrid-coordinate equations of motion have not yet found widespread application, but digital computer numerical integration programs have been applied to linear dynamic systems with nonlinear control at Hughes Aircraft Company (as described in Ref. 29), and similar programs are under development at the Jet Propulsion Laboratory.

Vehicle normal-mode coordinates have been employed in the simulation of controlled vehicles chiefly in application to missiles and launch vehicles. (Appropriate elastic-body equations may be found in Ref. 42.) Most applications of modal coordinates to spacecraft have been restricted to the determination of passive linear response of the spacecraft structure to its dynamic environment during launch. Both digital and analog computers have been employed successfully.

B. Linear System Analysis

The process of control system design often involves a preliminary phase in which it is assumed that the control system is to be linear, despite the designer's awareness that he will eventually devise a nonlinear control system. By assuming linearity at the outset, the designer gains access to simple analytical methods that he can employ quickly and efficiently to develop a preliminary control system design. Nonlinearities are then introduced into the system, and a detailed simulation is performed to confirm the acceptability of the design, or to provide the basis for its modification. A discussion of linear system analysis is included here for its utility in the preliminary design process.

Although many of the dynamic equations of the preceding sections have been nonlinear, it is frequently possible to replace each variable by the sum of a small variational coordinate and an explicit function of time that establishes the nominal controlled time behavior of that variable. (This was suggested in Sections III and IV, where \( a \) was replaced by \( \Theta(t) + \dot{\theta} \), with the variational coordinate \( \dot{\theta} \) presumed small.) In this way a linearized approximation can be obtained for any dynamic simulation of a controlled vehicle.

The linearized variational equations of the dynamic system can usually be combined with the linear equations of a preliminary control system design in a first-order equation as in

\[
\dot{Q} = BQ \tag{284}
\]

Here it has been assumed that it is possible to express the control torques as unknowns in a system of linear differential equations in the independent variable time, with coordinate coupling to the dynamic equations. In many cases the preliminary control torque will instead be available as an explicit function of the variational coordinates, in which case it can simply be substituted into the dynamic equations. Thus the matrix \( Q \) in Eq. (284) may or may not include control torques, but it will certainly include all of the kinematic coordinates and their time derivatives.

If the matrix \( B \) in Eq. (284) depends on time in an arbitrary way, nothing can be done with these equations but to integrate them numerically. In this event, the linearized equations are not substantially easier to deal with than their nonlinear counterparts.

If matrix \( B \) has a periodic time dependence, the stability of the null solution of Eq. (284) (corresponding to the
nominal motion) can be determined by application of Floquet theory. The equations would have this structure, for example, if for the dual-spin system simulated by the linear equations (224), (223), (222), and (216), the rotor were nonrigid or asymmetric. The application of Floquet theory to a similar problem may be found in Ref. 43.

In the simplest case, the matrix \( B \) in Eq. (284) is constant, and the stability of the null solution of this equation may be determined from the eigenvalues of \( B \). If the dimension of the matrix \( B \) is very small, or if Eq. (284) can be separated into a number of uncoupled matrix equations of small dimension, it may be practical to apply Routh's stability criteria (see any basic controls text, or Ref. 40, p. 9). Such an application is illustrated later in this section. In most cases of interest, however, the high dimension of matrix \( B \) necessitates the numerical (digital computer) calculation of eigenvalues.

The presence of any eigenvalue of \( B \) with a positive real part indicates the instability of the null solution of Eq. (284), and, by a basic theorem of Liapunov (see Ref. 34), also the instability of the null solution of the corresponding nonlinear equation.

If all eigenvalues of \( B \) have negative real parts, the null solution of Eq. (284) is asymptotically stable, as is that of the corresponding nonlinear equation.

If none of the eigenvalues of \( B \) has a positive real part, and one or more have a zero real part, no determination of the stability of the null solution of the nonlinear equation can be obtained from Eq. (284).

In many applications of interest, not every kinematic coordinate appearing in \( Q \) is relevant to the dynamic response of the system. For example, the coordinate \( \chi \), which describes the relative rotation of the symmetric rotor of the dual-spin spacecraft in Eqs. (224) and (223), does not appear (undifferentiated) in any of the dynamic equations of the system. Such a coordinate is termed cyclic or ignorable, and it must produce two zero eigenvalues for the matrix \( B \). Theoretically, one must then manipulate the differential equations so as to remove the cyclic-coordinate derivatives from Eq. (284), or re-derive the equations with the use of special methods that suppress these variables (see Ref. 24, p. 54). In practice, however, it may be sufficient simply to ignore those pairs of zero eigenvalues of \( B \) that can be identified with cyclic coordinates.

A digital computer eigenvalue program developed at Hughes Aircraft Company (noted in Ref. 29) has been used for dual-spin-satellite attitude-stability studies, based essentially on Eqs. (238), (223), (224), and (222) (with the last two equations written instead in terms of cantilever-appendage modal coordinates).

Preliminary design of linear control systems is traditionally based on the use of transfer functions, which establish the response of the Laplace transforms of the system variables to correspondingly transformed input torques. The hybrid-coordinate method is particularly compatible with this practice, although the method can be applied also when vehicle normal-mode coordinates are used.

Although Laplace transforms can be usefully applied to any system of linear equations, the advantages of this approach are most obvious when interest is focused on a small number of response variables. The concern of the attitude-controls engineer is generally limited to the rotational coordinates \( \theta_1, \theta_2, \theta_3 \), which have been used here to describe the small deviations of the base or main body of the spacecraft from its nominal attitude. Special problems are introduced when sensors or control actuators are located on flexible appendages, since then the appendage deformation coordinates enter the equation that establishes the control law. In any event, of course, the appendage vibrations must be permitted to influence \( \theta = [\theta_1, \theta_2, \theta_3]^T \), but often the appendage coordinates themselves are not of interest. With the Laplace transform approach, it is a simple matter to remove the (transformed) appendage coordinates from the (transformed) vehicle equations, and in this way to display most clearly the relationship between transformed control torque \( T(s) \) and transformed rotation \( \theta(s) \).

To illustrate this method without encumbrance, it will be applied here to the relatively simple problem of the inertially stabilized nongyroscopic space vehicle with a single flexible appendage. The appropriate dynamic equations may be obtained by specializing Eqs. (216) and (224) to exclude rotor and damper, and to eliminate external resultant force \( F \) and external torques \( \lambda \) on the appendages. The results are

\[
T = I^* \dot{\theta} + (\Sigma_{OE}^T + \vec{R} \Sigma_{EO}^T + \Sigma_{EO}^T \vec{r}) M \ddot{\vec{\rho}} \\
\dot{\vec{\rho}} + 2\xi \vec{\rho} \dot{\vec{\rho}} + \sigma^2 \vec{\rho} = -\vec{\rho}^T M (\Sigma_{OE} - \Sigma_{EO} \vec{R} - \vec{r} \Sigma_{EO}) \dot{\theta} \\
\text{(285)}
\]

\[
\text{Define now the } N \text{ by } 3 \text{ matrix} \\
\tilde{\delta} = -\vec{\rho}^T M (\Sigma_{OE} - \Sigma_{EO} \vec{R} - \vec{r} \Sigma_{EO}) \\
\text{(287)}
\]
and note from Eq. (142) that Eqs. (285) and (286) may be written as

\[ T = \Gamma^* \dot{q} - \delta^r \delta \eta \]  
\[ \ddot{\eta} + 2 \xi \sigma \dot{\eta} + \sigma^2 \eta = \delta \ddot{q} \]  

(The bar over \( \delta \) appears because the matrix of eigenvectors \( \tilde{\delta} \) has been truncated, but the definition of Eq. (287) is not so restricted.)

Equations (288) and (289) provide the following relationships among the Laplace transforms:

\[ \mathbf{D} = (\sigma^2 + 2 \xi \sigma \sigma + \sigma^2)^{-1} \]

When Eq. (292) is written in inverse form,

\[ \theta (s) = [s^2 \Gamma^* - s^4 \delta^r \mathbf{D} \delta]^T \Gamma (s) \]  

the matrix coefficient of \( T (s) \) is called the matrix of transfer functions. This matrix can be written in more convenient form in the special case in which truncation to a single modal coordinate has been imposed. In this case \( \mathbf{D} \) is a scalar, \( \delta \) is a 1 by 3 matrix (say, \( \delta^t \)), and Eq. (294) becomes

\[ \theta (s) = \frac{1}{s^2} \left[ \frac{1}{s^2 + 2 \xi \sigma \sigma + \sigma^2} \right]^{-1} T (s) \]  

The matrix \( \delta^r \delta^t \) (or \( \delta^r \delta^t \delta^r \) in this special case) is, from the definition in Eq. (278), a 3 by 3 matrix with the units of an inertia matrix. A physical interpretation of this matrix will be obtained indirectly in what follows.

It may first be noted that in the limiting case of a rigid appendage, with \( \sigma_i \) approaching infinity, Eqs. (294) and (295) provide the expected transfer functions for a rigid vehicle. The transfer-function matrix is then \( (\mathbf{I}^* - \delta^r) \mathbf{D} / s^2 \). If \( \mathbf{I}^* \) is diagonal, this is simply a collection of three single-axis transfer functions \( 1 / I_{z^\alpha}^2 \), \( \alpha = 1, 2, 3 \). Since \( \mathbf{I}^* \) is the inertia matrix of the entire vehicle about the vehicle mass center, these are the anticipated limiting-case transfer functions.

Consider now the opposite limiting case, for which the flexible appendages are so flexible that they are essentially detached, exerting no influence on the motion of the rigid body (i.e., all natural frequencies \( \sigma_i \) go to zero). Equations (294) and (295) then provide the limiting-case transfer functions from the expression

\[ \theta (s) = \frac{1}{s^2} (\mathbf{I}^* - \delta^r \delta^t)^{-1} T (s) \]  

This limiting case affords a physical interpretation of \( \delta^r \delta^t \) as the difference in inertia matrix \( \mathbf{I}^* \) of the total vehicle about its mass center and the inertia matrix \( \mathbf{I}^t \) of the rigid body to which the appendage is attached, referred to its own mass center. If \( \delta^t \) is the \( i \)th row of the matrix \( \delta \) defined in Eq. (287), and sufficient accuracy is preserved after truncation, the relationship

\[ \sum_{i=1}^{N} \delta^r \delta^t \delta^i = \mathbf{I}^* - \mathbf{I}^t \]  

must apply. In the special case in which the vehicle mass-center location is the same with or without flexible ap-
pendages, the matrix $\tilde{\delta}^T \tilde{\delta}$ is the matrix $I^4$ that appears in the lower right-hand corner of the $\delta$ by $6$ matrix $\Delta^T \Delta$ as expanded in Eq. (267).

For preliminary control system design, it is not unreasonable to consider the influence of a flexible appendage to be adequately represented by a single modal coordinate, permitting the use of Eq. (295) rather than the complex matrix expression found in Eq. (294). There then arises the question as to whether the matrix $\tilde{\delta}^T \tilde{\delta}$ in Eq. (295) should be replaced by its ideal value as in Eq. (297) or calculated from the definition of Eq. (287), truncating $\phi$ to a column matrix corresponding to the first mode shape. The use of the physical interpretation in Eq. (297) is of course simpler, since it does not require knowledge of the mode shape, but it is misleading in its implication that all portions of the appendage participate equally in the first modal vibration. The alternative of using the definition of $\tilde{\delta}$ from Eq. (287) does not give wholly accurate results either, however, for such severe truncation. In many situations, it is possible to combine experience and Eq. (297) to obtain as meaningful an estimate of the proper value of $\tilde{\delta}^T \tilde{\delta} = \delta^T \delta^T$ as the approximate transfer function of Eq. (295) warrants. Thus actual eigenvector (mode shape) calculations can often be avoided in very preliminary analysis.

The linearity of the system permits the use of superposition in generating from Eqs. (294) and (295) a more convenient transfer-function representation for use with truncation to $N$ modes, namely,

$$\theta (s) = \frac{1}{s^2} \left[ I^* - s^2 \sum_{i=1}^{N} \frac{\delta^T \delta^i}{s^2 + 2 \zeta_i \sigma_i s + \sigma_i^2} \right] T(s)$$

(298)

The traditional block diagram interpretation of a control system can very readily be imposed on flexible vehicles simulated with hybrid coordinates. Rather than simply substitute the transfer function of Eq. (294) or Eq. (298) into a single dynamics block $G(s)$, as shown in Fig. 12a, one can introduce a rigid-vehicle block $R(s)$ with feedback loops to accommodate the appendage flexibility corrections, as in Fig. 12b.

The detailed structure of the rigid-vehicle-block transfer function $R(s)$ and the flexibility-block transfer function $F(s)$ may be obtained by writing (with symbols from Fig. 12b)

$$\theta (s) = R(s) T^* (s) = R(s) [T(s) + F(s) \theta (s)]$$

(299)

or

$$[1 - R(s) F(s)] \theta (s) = R(s) T(s)$$

(300)

or

$$T(s) = [R^{-1}(s) - F(s)] \theta (s)$$

(301)
The transfer function $R(s)$ may be chosen so as to provide the rigid-vehicle term in Eq. (294), namely,

$$ R(s) = \frac{1}{s^2 (I^*)^{-1}} $$  

(302)

and the flexibility transfer function $F(s)$ then follows from Eq. (298) as

$$ F(s) = s^4 \sum_{i=1}^{N} \frac{\delta^i \delta^i}{(\delta^2 + 2\zeta_1 \sigma_1 s + \sigma_1^2)} $$  

(303)

An explicit representation of the breakdown of the dynamics block $G(s)$ to accommodate flexible appendages is shown in Fig. 12c.

Calculations similar to the preceding produce a transfer function for the total control system in the form

$$ T^R(s) = [G^{-1}(s) + H(s)] \theta(s) $$  

(304)

as may be verified by comparison with Eq. (301), making the necessary change of sign. The inverse of the matrix in brackets in Eq. (304) is the matrix transfer function of the system, and $G(s)$ is the inverse of the comparable matrix in Eq. (301). Equation (304) implies that the value of $\theta$ is sensed directly by sensors mounted on the rigid base.

If a sensor is mounted on a flexible appendage, it senses the sum of $\theta$ and any rotations due to structural deformation. Assume that the sensor is attached to the $i$th sub-body of the appendage, as portrayed in Fig. 4. Then $\beta^i$ is the 3 by 1 matrix of rotational deformations of the structure at the sensor location (see Eq. 72), so that $\beta^i$ is the $(2i)$th submatrix of dimension 3 by 1 in the deformation matrix $q$ (see Eq. 83). Recall that $q$ is related to $\eta$ by the transformation $q = \phi \bar{\eta}$, from Eq. (211). Thus $\beta^i$ may be obtained as the product with $\bar{\eta}$ of that 3 by $N$ portion of $\phi$ obtained as the $(2i)$th set of 3 by $N$ row partitions of $\phi$. Denote this 3 by $N$ matrix as $\phi_{2i}$, so that $\beta^i$ may be written as

$$ \beta^i = \phi_{2i} \bar{\eta} $$  

Thus the sensor reads $\theta + \phi_{2i} \bar{\eta}$ instead of $\theta$. The control system transfer function $H(s)$ would for a flexibly mounted sensor operate on $\theta(s) + \phi_{2i} \bar{\eta}(s)$, rather than on $\theta(s)$ alone. But Eq. (291) provides

$$ \bar{\eta}(s) = (s^2 E + 2s \bar{\zeta} \bar{\eta} + \bar{\sigma}^2)^{-1} s^2 \bar{\delta} \theta(s) $$

so that the control system transfer function $H(s)$ actually operates on $[E + \phi_{2i} (s^2 E + 2s \bar{\zeta} \bar{\eta} + \bar{\sigma}^2)^{-1} s^2 \bar{\delta}] \theta(s)$. Equation (304) is thus modified to the form

$$ T^R(s) = [G^{-1}(s) + H(s) (E + \phi_{2i} (s^2 E + 2s \bar{\zeta} \bar{\eta} + \bar{\sigma}^2)^{-1} s^2 \bar{\delta})] \theta(s) $$  

(305)

The dynamics transfer function $G(s)$ is then, from Eq. (301), the scalar

$$ G(s) = \left[ \frac{s^2 (\delta^i_0)^2}{(s^2 + 2\zeta_1 \sigma_1 s + \sigma_1^2)} \right]^{-1} $$

$$ = \frac{1}{(I_n s^2)} \left[ 1 - \frac{s^2 (\delta^i_0)^2}{I_n^a} \right]^{-1} $$

$$ = \frac{1}{(I_n s^2)} \left[ s^2 \left[ 1 - \frac{(\delta^i_0)^2}{I_n^a} \right] + 2\zeta_1 \sigma_1 s + \sigma_1^2 \right]^{-1} $$  

(307)

or

$$ G(s) = \left[ \frac{s^2 + 2\zeta_1 \sigma_1 s + \sigma_1^2}{(I_n s^2)} \left( s^2 \left[ 1 - \frac{(\delta^i_0)^2}{I_n^a} \right] + 2\zeta_1 \sigma_1 s + \sigma_1^2 \right) \right]^{-1} $$  

(308)
The quantity

\[ 1 - \left( \frac{\delta^2 \xi}{I_0} \right)^2 \]

which is 1 for a vanishingly small appendage and zero for an appendage on a vanishingly small base, is given the label \( R = 1 - \left( \frac{\delta^2 \xi}{I_0} \right)^2 \) (309)

and may be called the normalized reduced inertia for axis \( \alpha \) and mode 1.

On the assumption that the sensors are attached directly to the base, the total control system transfer function is available from Eq. (304) as the scalar

\[ \frac{\theta_\alpha(s)}{I_0 \theta_\alpha(s)} = \frac{G(s)}{1 + G(s)H(s)} \]  

(310)

From this expression, the customary procedures of control system synthesis can be applied. For example, stability can be determined (in the linear approximation) from the roots of the characteristic equation in \( s \),

\[ 1 + G(s)H(s) = 0 \]  

(311)

This is illustrated here for a simple gain control, so that \( H(s) \) is the constant \( K \), and Eq. (311) becomes (noting Eqs. 308 and 309)

\[ 1 + \frac{K(s^2 + 2\xi_1 \sigma_1 s + \sigma_1)}{I_0 \sigma_1^2 [R \sigma_1^2 + 2\xi_1 \sigma_1 s + \sigma_1]} = 0 \]

or, if \( R \sigma_1^2 + 2\xi_1 \sigma_1 s + \sigma_1 \neq 0 \), this becomes

\[ s^4 (I_0 \sigma) + s^3 \left( 2I_0 \xi_1 \sigma_1 \right) + s^2 \left( I_0 \xi_1^2 + K \right) + s \left( 2K \xi_1 \sigma_1 \right) + K \sigma_1 = 0 \]  

(312)

Stability is readily assessed for this system by the use of the Routhian array:

\[
\begin{array}{ccc}
\sigma_1: & I_0 \sigma & 1 \\
\xi_1: & I_0 \xi_1 & K \sigma_1 \\
\sigma_1: & 2I_0 \xi_1 \sigma_1 & \xi_1 \\
\xi_1: & I_0 \xi_1^2 + K - R \xi_1 & \sigma_1 \\
\sigma_1: & 2K \xi_1 \sigma_1 - \frac{2I_0 \xi_1 \sigma_1}{I_0 \sigma_1^2 + K - R \xi_1} & 0 \\
\sigma_1: & K \sigma_1 & 0 \\
\end{array}
\]

Necessary and sufficient criteria for asymptotic stability follow:

1. \( I_0 \sigma > 0 \)
2. \( 2I_0 \xi_1 \sigma_1 > 0 \)
3. \( I_0 \sigma_1^2 + K (1 - \sigma) > 0 \)
4. \( 2K \xi_1 \sigma_1 \left[ I_0 \sigma_1^2 + K (1 - \sigma) - I_0 \sigma_1 \right] = 2K^2 \xi_1 \sigma_1 (1 - \sigma) > 0 \)
5. \( K \sigma_1 > 0 \)

Assuming that gain \( K \) is positive, one may infer asymptotic stability in every case, since \( 0 < \sigma < 1 \) and all other symbols appearing in these criteria are positive.

Of course it should not be assumed that the appendage can never destroy control system stability; this supposition is demonstrated here only in a very special case.

It may be useful for the control system designer to construct root locus plots corresponding to simple models such as that reflected in Eq. (312), which is based on single-axis response of an appendage represented by a single modal coordinate. The root locus plot is simply the plot of the locus of the roots of Eq. (312) as \( K \) varies from zero to infinity. When \( K = 0 \), these roots (called the poles) are

\[ s_1 = p_1 = 0 \]
\[ s_2 = p_2 = 0 \]
\[ s_{3,4} = p_{3,4} = -\frac{I_0 \xi_1 \sigma_1 \pm \left[ (I_0 \xi_1 \sigma_1)^2 - I_0 \sigma \xi_1 \right]^{1/2}}{I_0 \sigma} \]
\[ = \left( \frac{\sigma_1}{\epsilon R} \right) [-\xi_1 \pm i(\epsilon - \xi_1)^{1/2}] \]  

(313)

and when \( K = \infty \), these roots (called the zeros) are

\[ s_{1,2} = z_{1,2} = \sigma_1 \left[ -\xi_1 \pm i \left( 1 - \xi_1 \right)^{1/2} \right] \]  

(314)

For most problems of practical interest, the percentage of critical damping is only a few percent, so \( \sigma - \xi_1 \) is positive. Even when this condition is violated, however, the poles in Eq. (313) remain in the left half-plane, as the Routh analysis guaranteed.
POLAR LOCI FOR VARYING $\zeta_1$ AND CONSTANT $\gamma$

FEATURES OF POLE LOCI FOR VARYING $\gamma$ AND CONSTANT $\zeta_1$:
- $\gamma = 1$: POLES coalesce at $-\sigma_1/\zeta_1$, 0
- $\gamma \to 0$: POLES diverge on real axis, approaching 0 and $\infty$

$\gamma = 0.17$, $\zeta_1 = 0.17$
$\gamma = 0.10$, $\zeta_1 = 0.10$
$\gamma = 0.05$, $\zeta_1 = 0.05$
$\gamma = 0$, $\zeta_1 = 0$

Fig. 13. Loci of poles and zeros
Figure 13 is not a root locus for $K$ ranging between zero and infinity, but is instead a plot of the loci of the poles and zeros as $\mathcal{R}$ and $\zeta_1$ vary. This figure is in the plane of solutions for $s = \alpha + i\beta$. It shows the locus of the two complex conjugate zeros as $\zeta_1$ varies from zero (when $z_{1,2} = \pm t\alpha$) to 1 (when $z_{1,2} = -1, -1$), and finally to infinity (when $z_{1,2} = 0, -\infty$). These loci are represented as long-dash lines in Fig. 13. The short-dash lines show the pole loci with $\zeta_1$ variation. The two poles at zero are unchanged, but the poles $p_3, p_4$ change from purely imaginary roots when $\zeta_1 = 0$ to double real (and negative) roots when $\zeta_1 = \mathcal{R}$. The poles then split, and approach zero and negative infinity as $\zeta_1$ increases. The zeros are uninfluenced by variations in $\mathcal{R}$, as are the two poles at zero. The two remaining poles coincide with and cancel the zeros when $\mathcal{R} = 1$, since there is then no flexible appendage on the vehicle. As $\mathcal{R}$ goes from 1 toward zero, the poles move along the solid-line loci in Fig. 13. When these loci reach the real axis (as shown for $\zeta_1 = 0.50$ and $\zeta_1 = 0.707$), the poles split and approach zero and negative infinity.

The more customary root locus plot shows the path of the roots from the poles (where $K = 0$) to a final position either at infinity or coincident with the zeros (where $K = \infty$). Such a plot requires repeated solution of the fourth-degree equation in Eq. (312), and is a suitable task for a computer. A few such plots have been generated for simple control system transfer functions $H(s)$, and no instabilities were found, although no systematic study has been undertaken. Figure 14 illustrates a typical result for the simple gain control system previously discussed, with $H(s) = K$. For such a system, it is a simple matter to establish the existence of vertical asymptotes, and to determine their intersection with the real axis. With a little additional labor, one can find general expressions for the lines of approach to the zeros and departure from the poles. Such calculations indicate that Fig. 14 is typical for practical values of $\mathcal{R}$ and $\zeta_1$. It is interesting to note that even as $\mathcal{R}$ approaches unity and the appendage poles and zeros approach coalescence, the loci that converge upon the zeros continue to be those rigid-body poles at the origin, as shown in Fig. 14, for values of $\mathcal{R}$ as high as 0.97.

The exploration of the influence of flexible appendages on control system transfer functions appears to be perhaps the easiest and most immediately rewarding path to understanding of system behavior, but this exploration has only begun.

Fig. 14. Root locus plot (variable gain $K$)
Nomenclature

\( \text{A} \) flexible appendage (generic symbol)

\( A' \) skew-symmetric matrix coefficient of \( q \) (see Eq. 140)

\( A_i \) \( i \)th flexible appendage

\( A_{ij} \) \( i \)th rigid sub-body of appendage \( A \)

\( a \) inertial acceleration of space vehicle mass center

\( a_i \) inertial acceleration of the \( i \)th body mass center

\( a_a \) inertial acceleration of space vehicle mass center

\( a_i \) inertial acceleration of the \( i \)th body mass center

\( A \) vector array \( (a_1, a_2, a_3) \)

\( B \) rigid body, space vehicle base

\( B_i \) \( i \)th rigid body

\( b \) scalar constant locating damper mass (see Eq. 108)

\( b_i \) dextral orthogonal unit vectors fixed in \( B \)

\( b \) vector array \( (b_1, b_2, b_3) \)

\( C \) direction cosine matrix relating \( \{a\} \) to \( \{b\} \) (see Eq. 34)

\( \text{CM} \) vehicle mass center

\( c \) vector from \( \text{CM} \) to point \( O \) fixed in \( B \) (see Eq. 36)

\( c \) matrix representation of \( c \) in basis \( \{b\} \)

\( D' \) symmetric velocity-coefficient matrix (damping matrix)

\( d \) dashpot constant (see Eq. 132)

\( E \) identity dyadic (see Eqs. 103, 125)

\( E \) identity matrix (see Eq. 56)

\( E^1, E^2, E^3 \) columns of 3 by 3 identity matrix

\( e, e \) position vector and corresponding matrix describing motion of \( \text{CM} \) in \( B \) due to moving parts (see Eq. 58)

\( F \) external force applied to vehicle

\( F^a \) external force applied to body \( A_i \)

\( F^b \) matrix representing \( F^a \) in basis \( \{a\} \) (see Eq. 55)

\( f' \) force applied to \( B_i \) by \( A \) (see Eqs. preceding 239)

\( f \) force applied to \( B_i \), excluding \( f' \) (see Eqs. preceding 239)

\( f', f \) matrices corresponding to \( f' \) and \( f \), in basis \( \{b\} \)

\( f_i \) matrix of forces on body \( A_i \), and top half of matrix \( \lambda \)

\( G' \) skew-symmetric velocity coefficient matrix (see Eq. 84)

\( G \) skew-symmetric matrix (see Eq. 141)

\( H \) vehicle angular momentum with respect to mass center

\( H^i \) angular momentum of body \( A_i \) with respect to its mass center

\( H^i \) matrix representation of \( H^i \) in vector basis fixed in \( A_i \)
Nomenclature (contd)

\( \mathbf{h}, \mathbf{h} \) angular momentum vector and \( \{b\} \) basis matrix for rotor \( B \), relative to base \( B \)

\( \mathbf{I} \) inertia dyadic for total vehicle with respect to point \( O \) (see Eq. 102)

\( \mathbf{I}' \) value of \( \mathbf{I} \) for undeformed vehicle (see Eq. 118)

\( \mathbf{I}' \) inertia dyadic of vehicle excluding appendage, with respect to \( P' \), the mass center of this portion of the vehicle (see Eq. 240)

\( \mathbf{I}^i \) inertia dyadic for \( A_i \) with respect to its mass center \( P_i \) (see Eq. 69)

\( \mathbf{I}, \mathbf{I}', \mathbf{I}' \) inertia matrices in basis \( \{b\} \) for dyadics \( \mathbf{I}, \mathbf{I}', \) and \( \mathbf{I}' \)

\( \mathbf{I}^i \) inertia matrix for \( \mathbf{I}^i \) in basis fixed in \( A_i \)

\( \mathbf{I}'^i \) inertia matrix of the undeformed appendage with respect to \( P' \) in basis \( \{b\} \)

\( \{i\} \) vector array of unit vectors \( i_1, i_2, i_3 \) fixed in inertial space

\( \mathbf{J} \) 6 by 6 expanded inertia matrix (see Eq. 258)

\( \mathbf{J} \) inertia dyadic of the rotor (see Eq. 107)

\( \mathbf{J} \) moment of inertia of rotor about symmetry axis

\( K' \) coefficient matrix for \( q \) (see Eqs. 84, 91)

\( k \) damper spring constant (see Eq. 132)

\( L' \) column matrix forcing function (see Eqs. 84, 94)

\( l' \) matrix of torques about mass center of body \( A_j \), and bottom half of matrix \( \lambda \)

\( \mathbf{l}', \mathbf{l} \) vector and \( \{b\} \) basis matrix for torque applied to \( B \), by \( A \) (see Eqs. 240 and 242)

\( l', l \) vector and \( \{b\} \) basis matrix for torque applied to \( B_i \), excluding \( \mathbf{l}' \) (see Eqs. 240 and 242)

\( M \) generalized 6n by 6n inertia matrix of cantilevered appendage (see Eq. 86)

\( M' \) generalized inertia matrix (see Eqs. 84 and 87)

\( m \) space vehicle total mass

\( m \) damper mass

\( m_i \) mass of \( i \)th body

\( m_i \) 3 by 3 mass matrix of \( i \)th body \( (m_i = m_i E) \)

\( m \) mass of system excluding appendage (see Eq. 239)

\( m' \) mass of appendage (see Eq. 267)

\( N \) either number of coordinates after truncation (as in Eq. 166), or number of total coordinates in a constrained set (as in Eq. 22)

\( N' \) nonlinear terms in Eq. (140)

\( n \) either total number of rigid bodies in a discrete parameter model of an appendage (as in Fig. 4), or total number of rigid bodies in a discrete parameter model of a vehicle

\( O \) point fixed in \( B \), coincident with \( CM \) when vehicle undeformed
Nomenclature (contd)

\( O' \) point fixed in inertial space (see Eq. 35)
\( P \) 12\( n \) by 12\( n \) transformation matrix (see Eq. 179)
\( P' \) mass center of vehicle exclusive of appendages
\( p \) either generic position vector from \( O \) (see Eq. 98), or vector from \( P' \) to \( Q \) (as in Eq. 246)
\( p^* \) value of \( p \) when vehicle undeformed
\( p^A \) matrix in basis \{a\} of vector from \( P' \) to appendage mass center
\( P_s \) mass center of body \( A \),
\( Q \) either point fixed in \( B \) on interface of \( A \) and \( B \) (as in Eq. 36), or 12\( n \) by 1 \( n \) state variable (as in Eqs. 144, 145)
\( Q_s \) mass center of \( A \), when appendage undeformed
\( q \) 6\( n \) by 1 matrix of deformation coordinates (see Eq. 83)
\( R \) vector from \( O \) to \( Q \)
\( R \) 3 by 1 matrix representing \( R \) in basis \{b\}
\( r, r_s \) vector from \( Q \) to \( Q_s \), and corresponding matrix in basis \{a\}
\( s \) Laplace transform variable
\( T \) 3 by 1 matrix of applied torque (see Eq. 128)
\( T_i \) torque applied to \( i \)th body about its mass center
\( t \) time
\( U \) 6 by 1 matrix of rigid-body coordinates (see Eq. 257)
\( u^*, u_s \) vector from \( Q_s \) to \( P_s \) and corresponding matrix in basis \{a\}, defining translational deformation of appendage (see Fig. 4)
\( X, X \) vehicle mass center inertial position vector and corresponding inertial basis matrix
\( x, x \) inertial position vector of point \( O \) and corresponding inertial basis matrix
\( Y \) transformed state variable (see Eq. 159)
\( Z \) transformed state variable (see Eq. 187)
\( Z^A \) 12\( n \) by 1 matrix of homogeneous solutions (see Eq. 177)
\( z \) 6\( n \) by 1 matrix of variables (see Eq. 186)
\( z_i \) elements of \( z \)
\( \sigma_k \) real part of \( \lambda_k \) (see Eq. 164)
\( \beta^* \) rotation vector for small rotations of body \( A \), relative to \( A \)
\( \beta^* \) matrix representation of \( \beta^* \) in vector basis fixed in \( A \)
\( \beta^+_1, \beta^+_2, \beta^+_3 \) elements of \( \beta^* \)
\( \Gamma^K \) imaginary part of \( \phi^K \), a 12\( n \) by 1 matrix (see Eq. 164)
\( \gamma^m \) upper half of \( \Gamma^m \) (see Eq. 174)
\( \gamma \) 6\( n \) by 6\( n \) matrix (see Eq. 186)
\( \Delta \) 6\( n \) by 6 matrix (see Eq. 261)
Nomenclature (contd)

\[ \Delta^T \Delta \] matrix product with interpretation as an inertia matrix (see Eq. 267)

\( \delta \) 6n by 3 matrix

\( \delta \) N by 3 matrix formed by truncating \( \delta \) (see Eq. 257)

\( \delta^i \) 1 by 3 matrix formed by truncating \( \delta \) to \( \delta \), with \( N = 1 \)

\( \zeta_m \) percentage of critical damping in vibration mode \( m \)

\( \zeta \) diagonal matrix with diagonal elements \( \zeta_1, \ldots, \zeta_n \)

\( \eta \) 6n by 1 matrix of modal coordinates (see Eq. 211)

\( \eta^c \) 6n by 1 matrix of modal coordinates for cantilever modes (see Eq. 232)

\( \Theta \) direction cosine matrix establishing inertial orientation of body \( B \) (see Eq. 53)

\( \hat{\theta} \) 3 by 1 matrix with elements \( \hat{\theta}_1, \hat{\theta}_2, \hat{\theta}_3 \), providing a linear approximation of \( \psi \) (see Eq. 130)

\( \Lambda \) 6 by 1 matrix of external forces and torques (see Eq. 258)

\( \lambda \) 6n by 1 matrix of external forces and torques (see Eq. 94)

\( \lambda_m \) eigenvalue (see Eq. 162)

\( \mu^* \) mass ratio \( m^*/m \)

\( \nu \) number of constraint equations

\( \xi \) translation of damper mass (scalar)

\( \rho \) generic position vector from vehicle mass center (see Eq. 97)

\( \rho^* \) generic position vector from mass center \( P_s \) of sub-body \( A_s \) (see Eq. 109)

\( \Sigma_{EO}, \Sigma_{OE} \) (see list of operators)

\( \sigma_{k} \) imaginary part of eigenvalue \( \lambda_s \), and hence, a natural frequency (see Eq. 164)

\( \sigma \) diagonal matrix with diagonal elements \( \sigma_1, \ldots, \sigma_{6n} \)

\( \tau \) rotor torque magnitude (see Eq. 136)

\( \Phi^m \) 12n by 1 eigenvector matrix (see Eqs. 147 and 148)

\( \Phi \) 12n by 12n transformation matrix with columns \( \phi^1, \ldots, \phi^{12n} \) (see Eq. 153)

\( \phi \) rotor angular speed (see Eq. 135)

\( \phi^c \) 6n by 6n transformation matrix for cantilever modes (see Eq. 232)

\( \phi^m \) 6n by 1 matrix describing mode shape (see Eq. 148)

\( \phi^m_i \) scalar \( i \)th element of matrix \( \phi^m \)

\( \phi \) 6n by 6n transformation matrix with columns \( \phi^1, \ldots, \phi^{6n} \) (see Eq. 210)

\( \Psi^K \) 12n by 1 matrix, the imaginary part of \( \Phi^K \) (see Eq. 164)

\( \psi^K \) 6n by 1 matrix, the imaginary part of \( \phi^K \), and upper half of \( \Psi^K \) (see Eq. 174)

\( \Omega^a \) angular velocity of \( \{a\} \) relative to \( \{b\} \) (see Eq. 48)

\( \Omega^a \) matrix representation of \( \Omega^a \) in \( \{a\} \) basis
Nomenclature (contd)

\[ \Omega \] angular velocity of rotor \( B_3 \) relative to body \( B_1 \) (see Eq. 107)
\[ \omega \] inertial angular velocity of \( B \)
\[ \omega^i \] inertial angular velocity of body \( A_i \)
\[ \omega^1, \omega^2, \omega^3 \] scalar measure numbers of \( \omega^i \) in basis fixed in \( A_i \)

Operators

\[ [\ ] \] square brackets enclose matrices only when the elements of the matrix are recorded explicitly (see Eq. 44)
\[ \{ \} \] braces always enclose vector arrays, which in this report are always column arrays of three orthogonal unit vectors (see Eq. 33)
\[ \sim \] tilde either over a symbol or as a superscript, denotes a skew-symmetric matrix formed from a column matrix, according to the pattern established by either Eq. (44) or Eq. (89), depending on the matrix dimension
\[ \cdot \] dot over a vector or dyadic denotes time differentiation in an inertial frame of reference. Dot over a scalar or matrix denotes time differentiation
\[ \circ \] circle over a vector or dyadic denotes time differentiation in the reference frame of body \( B_1 \) (see Eq. 98)
\[ \frac{d}{dt} ( ) \] denotes time differentiation in reference frame \( f \). (The symbol \( f \) may be replaced by \( a, b, \) or \( i \), which denote reference frames fixed respectively in body \( A \), body \( B \), or inertial space.)
\[ \sim \] bar over a matrix indicates truncation, changing dimension \( 6n \) to \( N \). (Note that \( \omega^i \) in Eq. 283 is excluded)
\[ \Sigma \] summation operator
\[ \Sigma_0, \Sigma_{0e} \] summation matrices consisting of ones and zeros (see Eq. 85)

Superscripts

\[ T \] denotes matrix transpose
\[ -1 \] denotes matrix inverse
\[ * \] as superscript for a complex number, denotes the complex conjugate

References

References (contd)


References (contd)


References (contd)


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