Implementation of topology optimization using openMDAO

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Topology optimization is one of the widely known branches among the structural optimization, and it distinguishes itself by a large design domain and versatility. It can determine the optimal design out of an infinite number of configurations, thereby drawing interest from both industry and academia with regards to its applicability to additive manufacturing. However, its implementation is often a daunting task for engineers in practice. One of the issues is the programming effort required whenever the implementation requires any changes, ranging from subtle tweaks to drastic changes in the problem definition, and the derivative computation must be correspondingly updated after any change. The implementation, therefore, is not only time-consuming but also repetitive and susceptible to human-induced errors. In this regard, topology optimization implementations stand to benefit from reusability, ease of restructuring, and modularity. In this work, we propose OpenMDAO, a computational framework for multidisciplinary design optimization (MDO), as a generic platform for topology optimization. Two widely used topology optimization techniques—density-based and level-set—are implemented as a demonstration. These techniques are implemented in a decomposed manner, with the aid of the modular architecture of OpenMDAO as well as state-of-the-art numerical methods. Variations on the density-based topology optimization approach are shown to demonstrate the modularity and automation for derivative computation that OpenMDAO provides.

Nomenclature

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Topology optimization is a numerical design method that computes an optimal structure for a specified objective and constraint. The method is capable of handling a large number of design variables, and thus can compute the optimal structural configuration from a large design space with minimal designer input.

Topology optimization methods can be classified into two categories based on the representation of the structural topology. One is the density-based formulation where the problem is formulated as a material distribution problem and the material densities are design variables. Typically, each element in the finite element mesh is separately parameterized as in Solid Isotropic Material with Penalization (SIMP) \cite{SIMP} method, which is the most common approach. Another category is the boundary-based formulation where the structure is defined by its boundary. The change of the structural topology is realized by the direct movement of the boundary encompassing merge, split, and creation operations. Its history can be traced back to shape optimization \cite{shape_optimization1, shape_optimization2}, which is inspired by computational geometry \cite{computational_geometry}. The introduction of the level-set method \cite{level_set1, level_set2} attracted much attention in recent years and is quickly becoming a viable alternative to SIMP due to its lack of a need for post-processing. Both methods employ the finite element method (FEM) as the backbone that computes the responses of the structure and the corresponding sensitivities with respect to the design variables. Since FEM can solve diverse partial differential equations that range from elasticity \cite{FEM_elasticity, FEM_acoustics} to acoustics \cite{FEM_acoustics}, it is also possible to perform multidisciplinary topology optimization. In recent years, topology optimization has been actively studied with Additive Manufacturing (AM) in mind, a novel manufacturing technique with far greater design freedom when compared to the classical methods. AM offers small-scale manufacturing capability with micron-scale resolution and multiple materials without additional costs, and leads to inherently multi-scale and multi-functional structures.

In practice topology optimization implementations can be somewhat difficult to implement. One strategy to simplify the implementation is the use of object-oriented programming (OOP) to create a modular code base with a high degree of code reuse. This approach also helps a user easily change the constituent units of code and their organization within the program. These OOP traits reduce the repetitive programming tasks that are not only time-consuming but also make
the code susceptible to human error during the implementation.

We propose OpenMDAO, a computing framework for Multidisciplinary Design Optimization (MDO), as a generic platform for topology optimization. This enables topology optimization to access a wide range of MDO methods for coupling multiple disciplines, and it facilitates integration of multiple computational tools. First, OpenMDAO uses a modular software architecture for the MDO problem, by which the model is configured into smaller computational units that communicate to each other, and OpenMDAO automatically computes the total derivatives of the model from those of the smaller units. Such an architecture is beneficial for restructurability with minimal effort. Moreover, OpenMDAO is equipped with a large library of nonlinear and linear solvers as well as visualization utilities. Based on these attributes, potential users can easily implement and reconfigure their optimization workflow, and easily share their modules and ideas.

These properties are also beneficial to researchers who are new to topology optimization. Once the framework is separated into smaller units of code and robustly programmed, external users can freely rearrange and tweak these components. These traits distinguish the present work from the previous educational publications on topology optimization [9, 10].

The first part of this paper defines the optimization problem and briefly describes the two topology optimization schemes implemented in this work: SIMP topology optimization and level-set topology optimization (LSTO). For both, we consider a linearly elastic compliance minimization problem. In the next section, the OpenMDAO implementations of SIMP and LSTO are explained with design structure matrix diagrams, and numerical examples are presented. The last part of the paper demonstrates an extension of the SIMP implementation to a third topology optimization formulation, the parametric level set approach. This extension provides a practical demonstration of the benefits of modularity, as we reuse existing components from the SIMP approach, add new components, and use OpenMDAO to integrate them and compute their derivatives with minimal effort.

II. Theory

A. Optimization problem

Compliance minimization is a classical problem in topology optimization [1], and the optimization problem is given by

$$
\min f(x, u) = u(x)^T F = \int_{\Omega} \nabla u(x) : E(x) : \nabla u(x) d\Omega
$$

$$
s.t. \quad g(x) \leq g^*
$$

(1)

where $\Omega$ is the domain, $f$ is the compliance of the structure, which is determined by design variable $x$ and displacement function $u$. $g(x)$ is the material volume found within the structure, $g^*$ is the prescribed volume constraint, and Young’s modulus $E$. Note that $u$ is also a function of $x$ because linear elasticity as a boundary value problem is assumed herein.
In this work, we look at a 2D cantilevered plate for demonstrating the implementations of both topology optimization methods. The design domain, boundary conditions, and the external loading are illustrated in Figure 1.

![Diagram](image)

**W = 2, H = 1**

**E = 1, v = 0.3**

**Fig. 1** The cantilevered plate for the compliance minimization problem. The dimensions of the design domain (width \( W \) and height \( H \)), Young’s modulus \( (E) \) with Poisson ratio \( (v) \), and external force \( (F) \) are illustrated.

In this work, the length-to-width ratio \( (W/H) \) is fixed to 2, and the Young’s modulus \( E \) and Poisson ratio \( v \) are set to 1.0 and 0.3, respectively. Force \( F \) is applied in the \(-y\) direction with a value of 1 at the mid-point of the right edge. A linear elasticity assumption is employed, and the computation of the design sensitivity \( \frac{dF}{dx} \) exploits the self-adjoint characteristic of the compliance minimization problem. The constraint \( g(x) \), the ratio of the total amount of material over the area of the whole domain, is set to be 40% (i.e. \( g^* = 0.4 \)). These optimization settings are fixed throughout this work to simplify the verification of the numerical results and the demonstration of the reusability and the restructurability of the current implementation. Note that design variable \( x \) that represent the structure varies depending on the type of topology optimization method. In SIMP, discrete material density \( \rho \) is used and level-set method utilizes level set function \( \phi \).

**B. SIMP: Density-based method**

In SIMP, the topology of the structure is explicitly described by the distribution of the discrete element-wise material density \( \rho \) that parameterizes the material property. The material densities \( \rho \) for each element are the design variables of SIMP, and therefore the number of design variables coincides with the number of elements. Due to its simplicity of implementation, the SIMP method has been widely used \[9,11\] for topology optimization. Without losing generality, Eq. (1) can be reformulated as follows

\[
\begin{align*}
\min c(\rho, u) &= \sum_{e} N_e \rho_e u_e^T k_e^\rho u_e \\
\text{s.t.} \quad &\sum_{e} \rho_e \leq G^* \\
\text{s.t.} \quad &K(\rho)u = F \\
\text{s.t.} \quad &0 \leq \rho_e \leq 1
\end{align*}
\]

(2)
where $p$ is a penalizing parameter, $u_e$ and $\rho_e$ are the its discretized displacement and material density correspond to element $e$, and $k^0_e$ is the stiffness matrix, provided that the element is filled with material (i.e. $\rho_e = 1$). A typical workflow is described in many works\cite{1, 9}, and it is illustrated in Figure 2.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{xdsm.png}
\caption{An XDSM diagram of SIMP method}
\end{figure}

The simplest decomposition of SIMP consists of 5 modules: filter, penalization, finite element analysis, objective, and constraint. The finite element analysis is a computational backbone of the SIMP method, as all inputs and outputs of each module are present in the same discretized space. Whenever the material connectivity is required in the problem definition, either density or sensitivity filter is required. In this diagram, the density filter is shown.

As shown in the diagram, the first step in the SIMP method is an applying density filter to the material distribution $\rho_i$ to generate locally uniform distribution $\bar{\rho}_i$. Then a filtered material density is penalized so that intermediate densities (i.e. gray material) are removed. The penalization is a crucial step in obtaining a well-defined topological solution. Without it, a converged solution is likely to contain gray areas that are open to interpretation during post-processing. In this work, the penalization parameter $p$ is fixed to 3. The material properties (i.e., Young’s modulus) are assumed to be uniform within each element, and proportional to the specific material densities found on the element as

$$E = \rho^p E_0 + (1 - \rho^p)E^\epsilon$$  \hspace{1cm} (3)$$

where $E_0$ is the original modulus of the material and $E^\epsilon = 10^{-6} E_0$ is a fictitious stiffness, which is introduced to prevent rank deficiency of the stiffness matrix and enforce nonzero sensitivity at the element. The objective and constraint functions are computed based on these variables.

Recalling that gradient-based optimization is utilized in this work, the update of the solution requires the derivatives of the objective function with respect to design variables. As there is a substantially higher number of design variables
than governing equations, the adjoint method is employed for efficiency. Due to the self-adjoint characteristic of the compliance minimization problem, the total derivative calculation is greatly simplified. The corresponding sensitivity to the material density is

\[
\frac{dc(\rho,u)}{d\rho_e} = -(p_f e_p^{-1}) u^T_{e} K_e(\rho) u_e
\]

where \( \rho \) is a material density, \( u \) is a displacement vector, and \( K \) is a stiffness matrix. The subscript \( e \) refers the element index found in the finite element mesh.

C. LSTO: level-set topology optimization

Level-set topology optimization (LSTO) differs from SIMP, as the structural domain is implicitly represented. An advantage of LSTO over density-based optimization scheme is the resulting boundaries of the optimal solution can cut an element and smooth boundaries are obtained in contrast to jagged-edge boundaries found in SIMP. In this work, we implemented the LSTO method found in the reference [12].

For a given level set function \( \phi(r) \) in the fixed grid space, the boundary is defined by its zero hypersurfaces \( (\Gamma(r) : \{ r \in \Omega | \phi(r) = 0 \}) \). A signed distance function, a distance measure from the boundaries of the design domain, is commonly used as a level set function, and it is implicitly updated by solving the Hamilton-Jacobi equation

\[
\dot{\phi} + \nabla \phi \cdot \frac{\partial r}{\partial t} = \phi + V_n |\nabla \phi| = 0
\]

where \( t \) is the pseudo-time in which the level set function advects, and \( V_n \) is the velocity of advection, normal to the structural boundary. An explicit integration method is often employed to solve the problem for the discretized space and time domain. Equation (5) is linearized with respect to shape perturbations (i.e., shape derivatives) at the boundary points.

\[
\begin{align*}
\min \frac{\partial f(x)}{\partial \Omega^k} \cdot \Delta \Omega^k &= \Delta t \int \Gamma S_f V_n d\Gamma \\
\text{s.t.} \frac{\partial g}{\partial \Omega^k} \cdot \Delta \Omega^k &= \Delta t \int \Gamma S_g V_n d\Gamma \leq -g^{*k}
\end{align*}
\]

where \( \Delta \Omega^k \) is the update for the current domain at the \( k \)-th pseudo timestep, and \( S_f \) and \( S_G \) each represent the shape sensitivities for the objective and the constraint. The function change is now a function of the shape sensitivities and normal velocities evaluated using boundary integrals.

In contrast to SIMP, the shape sensitivities are evaluated at the discretized boundary points, and the consistency and the smoothness of the values are necessary to obtain the optimal solution with clear distinction of the boundary. The consistent compliance sensitivities [6] are...
\begin{align*}
    S_f &= -\int_F \varepsilon(u) \cdot E(\rho) \cdot \varepsilon(u) dl \\
    S_g &= -1
\end{align*}

where the weighted least squares method is employed based on sensitivities \( S_f \) and \( S_g \) evaluated at the point clouds. The Gauss quadrature points are selected as the point of evaluation, and an Ersatz material model (i.e., \( E = E_0 \rho \), where \( \rho \) is an area fraction that is a ratio of materials presence \( (\phi > 0) \) within each element) is employed. It is worth noting that the sensitivity found in Eq. (7) can be reduced to Eq. (4) when the sensitivity is evaluated at the midpoint of each element as it also exploits the self-adjoint trait of the given problem.

As suggested by Dunning et al., \cite{7}, the linearized equation (6) leads to Eq. (8) after the algebraic manipulations. Detailed procedures also can be found in the recent literature \cite{12}.

\begin{align*}
    \frac{\partial [f, g]}{\partial \Omega^k} \cdot \Omega^k &= \sum_{n_b} \Delta t V_{nj} [S_{f,j}, S_{g,j}] l_j \\
    &= [C_f, C_g] \cdot V_n \Delta t \equiv [C_f, C_g] \cdot z \\
    z(\lambda) &= \lambda_f S_f + \lambda_g S_g
\end{align*}

where subscript \( j \) indicates the index of the boundary points, and \( C_f \) and \( C_g \) refer the vectors obtained by boundary integration of the corresponding sensitivities, while \( l \) refers the length segment. Normal velocity \( V_n \) is multiplied by the \( t \) to produce adveacting distance \( z \) that the boundary advects during given time, while \( z \) is assumed to be the sensitivities multiplied by Lagrange multiplier \( \lambda \). By using Eq. (8), Eq. (6) is then written as:

\begin{align*}
    \min_C f^T z \\
    s.t. \quad C_g \cdot z \leq g^* \\
    s.t. \quad z_{\min} \leq z \leq z_{\max}
\end{align*}

where a bound for a distance vector \([z_{\min}, z_{\max}]\) satisfies two criteria: (1) the advection must not go beyond the boundary of the design domain, and (2) the distance cannot violate CFL (Courant-Frederichs-Lowy) condition. These set of equations leads to the sub-optimization of the LSTO method, by which optimal distance for the function minimization is obtained and remains within the range of constraints and bounds.

As a result, two sets of the optimization layers are separately dedicated to the Hamilton-Jacobi equation and the sub-optimization, and shown in the XDSM diagram shown in Figure 3.
For a given signed distance function $\phi$, the boundary points are obtained by using a marching square algorithm\[4\]. The positions of the boundary points and the length segments of the boundary $l$ that represent discretized design and the resulting boundary sensitivities $S_f$ and $S_g$ are the arguments of the sub-optimization.

In order to retain distance $z$ within its bounds, the bounds of the multipliers $\lambda$ are also computed based on the CFL condition and the geometric limit of the movement (i.e., side limit \[13\]). These are the conservative move limits for boundary advection. The sub-optimization problem (Eq. \[9\]) is then solved and the optimal distance $z$ is computed as its solution. These boundary distances are extended to level-set fixed grid within the narrow band\[4\], and the structure is updated by solving the Hamilton-Jacobi equation. Furthermore, reinitialization of the function is required to satisfy the $|\nabla \phi| = 1$ condition as it is often violated after a few updates.

Note that the solution update scheme in LSTO requires time integration over pseudo time $t$, and involves sophisticated numerical techniques such as the fast-marching method. Due to these traits, the first layer of the optimization found in the LSTO is not benefited from the numerical solvers found in OpenMDAO. In this work, therefore, only the sub-optimization step is exposed to the OpenMDAO, where the optimal advection velocity at each optimization steps is computed. The outer part of the optimization, on the other hand, is solved externally by using numerical libraries for a level set, such as the upwind scheme, fast marching methods, and the 5th order WENO \[12, 14\].

### III. Implementations of topology optimization methods

#### A. OpenMDAO

OpenMDAO \[15\] is an MDO platform developed and maintained by researchers at NASA Glenn Research Center. OpenMDAO is open-source and is written in Python, which is a language known for its ease of interfacing to compiled languages such as C++ and Fortran. OpenMDAO has been used to solve MDO problems in satellite design \[16\], wind
turbine design [15], aircraft design [17], and aircraft trajectory optimization [18].

OpenMDAO is unique among MDO frameworks because it is designed for gradient-based optimization, and assists in the computation of derivatives. It uses the modular analysis and unified derivatives (MAUD) architecture [19] to compute the global derivatives of the model given the partial derivatives of each smaller unit of code. MAUD is a generalization of the adjoint method that unifies all existing methods for computing derivatives using a matrix equation [20]. Like the adjoint method, MAUD has the property that the computational cost of computing a gradient is independent of the number of design variables. MAUD uses parallel hierarchical solvers for solving the systems of equations that arise in coupled simulations.

![Illustration of modularized architecture found in OpenMDAO. Three set of the layers are found at the different hierarchies: Component, Group, and Problem.](image)

The basic unit of code in an OpenMDAO model is the Component, where the computations take place during an execution of OpenMDAO. It can be as comprehensive as a single discipline, but more often, it is a small function such as the penalization equation found in the SIMP-based optimization. The Component object is defined by its variables (inputs and outputs), the governing equation that computes the outputs for given inputs, and the partial derivatives that are either specified by the user or approximated by the OpenMDAO using finite differences or the complex-step method. Depending on the type of equation, a Component can be of either explicit or implicit type. For instance, a penalization module is an explicit Component, whereas the computation of the displacement vector using linear elasticity is an implicit one as the displacement $u$ is a state variable defined implicitly by residual functions. In the latter example, $u$ is nonetheless an output variable that the Component is responsible for computing.

A Group is located above the Component objects in the model hierarchy tree and contains a combination of Component objects and other Group objects. It is a common practice to assign each individual discipline its own Group,
and its algorithms are realized by placing Components within the Group and arranging them in a logical order.

The configuration of the Group object is greatly simplified thanks to the MAUD architecture in OpenMDAO. First of all, there is a salient modularity found during the process due to the encapsulation of the Component object; only the arguments of the Components are exposed to the Group layer, while the others are remain hidden. Moreover, the total derivative is automatically computed as long as the partial derivatives of the constituting Components are specified. Such capability is especially useful in the presence of the implicit Component as the adjoint method is internally evoked in OpenMDAO without resorting to the explicit calculation.

Such a modular structure leads to both restructurability and reusability. Once the Component object is programmed and encapsulated, a further modification is unnecessary as long as the governing equations remain intact (reusability). Also, these modules are freely arranged within the Group layer and therefore reformatting of the information requires only changes in connectivity between the Components, and the required changes to the global derivative are applied automatically (easy restructurability). These two traits are beneficial as these significantly reduce repetitive programming and user-induced error.

The model is thus a combination of Group and Component objects, arranged in a hierarchy tree. The leaves of this tree are comprised of Component objects and all other nodes in the tree are Group objects. The model is contained within the Problem object, which also contains a Driver object. The Driver is responsible for execution of the model and for computation of the model’s derivatives using MAUD. It is typically the optimization algorithm. For the present study, the Driver is an SLSQP optimizer with a tolerance of $10^{-9}$.

In this work, topology optimization is deemed as the one discipline of our MDO problem, and a single Group object is therefore assigned to each optimization method (e.g., SIMP, LSTO). In our case, there are no other Group objects in the model. Therefore, each Group is comprised solely of Component objects, including the finite element analysis and auxiliary computations such as the density filter found in the SIMP method. Figures 2 and 3 explain the topological optimization workflows visually using design structure matrix and hierarchy diagrams. Each Component is marked by the blue box, and its inputs (pink-colored) and outputs (either gray- or orange-colored) exposed to Group layer are also specified. An orange-colored output is an unknown from implicit function, while gray-colored one is an explicit output. Note that dependencies are marked by black off-diagonal squares.

**B. SIMP**

The visualization of SIMP is shown in Figure 5, where the Components and their connectivity in the SIMP Group are illustrated.

The workflow of the SIMP method is composed of six Components: specifying independent variables (input_comp), density filtering (filter_comp), penalization (penalization_comp), linear elasticity (states_comp), the objective function (compliance_comp), and the constraint function (weight_comp). Note that this configuration is analogous to
the modules and their connectivities found in the corresponding XDSM diagram (Figure 2). For example, displacement (disp) and forces (forces) are inputs to the compliance_comp, of which output is a compliance.

After preprocessing that includes discretization and specifying boundary conditions, the independent inputs are specified for the given finite element mesh: discrete material density \( \rho_s \) as a design variable and external force vector forces. Design variables first go through a simple density filter, a low-pass filter that removes a checkerboard pattern often found within the solution:

\[
\bar{\rho}_i = \sum_{j} w_{ij} \rho_j, \quad w_{ij} = \frac{R - d(i, j)}{\sum_{k \in N_i} (R - d(i, k))} \quad \text{if } j \in N_i
\]

where \( \bar{\rho}_i \) is a filtered material density found at the element \( i \), and \( w_{ij} \) is a conic weight based on the distance \( d \) between elements \( i \) and \( j \). A Group \( N_i \) refers to the list of the neighboring elements of element \( i \), within a radius \( R \). Although the filtering is not visible in describing XDSM of SIMP method (Figure 3), it is commonly employed for a stable convergence in SIMP. Choosing the best filter, however, relies heavily on the heuristics; therefore, iterative testing is often required [21]. Again, the restructurability of the present approach reduces the effort required.

The filtered design variables are then penalized with constant \( p \). The computed material density \( \bar{\rho}_p \) (multipliers) leads to parameterized material stiffness (Eq. 3) and global stiffness matrix. The displacement vector states is a state variable by solving force equivalence \( (R(\rho, u) = K(\rho)u - F = 0) \). Within the state_comp Component, the partial derivatives of the residual \( R \) with respect to all inputs are computed, which are later used by OpenMDAO in the adjoint
method. This is another advantage of using OpenMDAO as a framework of topology optimization as it greatly simplifies the calculation of the global gradient.

![Numerical results of SIMP](image)

**Fig. 6** Numerical results of SIMP. (a) a convergence profile with initial configuration where the number of meshes to each direction are specified (b) a material configuration, area fraction ($g$) and compliance ($f$) found during at the 4th, 45th, and 200th iterations.

Figure 6(a) shows the convergence graph of the compliance $f(\rho)$, which is a objective function, the area fraction $g$, and the initial configuration of the design domain. The size of the rectangular design domain is prescribed to 160 x 80 (length x width), and the domain is discretized with 80 x 40 quadratic elements. The initial material densities are uniformly set to 0.5, by which the constraint is mildly violated. According to the convergence graph, a constraint ($g(\rho) < 0.4$) is always satisfied after 5th iteration and the norm of the changes of the compliance is smaller than $10^{-6}$ after 48th iteration. However, the iteration continues until the 200th iteration because of the small convergence criteria ($\Delta f(\rho, u) < 10^{-9}$). In Figure 6(b), the final structural topology is nearly obtained. Note that intermediate densities found in the early iterations are penalized so that the solutions at the later iterations do not exhibit the intermediate densities.

C. LSTO

As described in its XDSM diagram (Figure 3), LSTO is composed of two layers: (1) an update of the level-set function achieved by solving on Hamilton-Jacobi equation on the domain boundary, and (2) a sub-optimization, by which the optimal advection distance at each iteration is calculated. As mentioned earlier, only the sub-optimization step in Eq. (9) is implemented within OpenMDAO. On the other hand, the pre-processing operations and the post-processing operations, such as updating and extracting the design, are separately accomplished outside of OpenMDAO.

The operations prior to sub-optimization consist of initializing the level-set properties and execution of the finite
element analysis. A signed distance function is firstly initialized, and the geometric properties of the structure, which includes the location of the boundary points, length segments, and side limits of the boundary movement, are extracted by external level-set libraries based on Ref. [13]. The structure in the finite element domain is represented by the Ersatz material model, and the boundary conditions are assumed to be constant during the iteration. As a result, the sensitivities $S_f$ and $S_g$ evaluated at the boundary points and the geometric properties are obtained, based on which the optimal distances at the structural boundaries are calculated through sub-optimization. The post-processing operations update the level set function based on the calculated distance. The given distances $z$ at the boundary points are first extended to the level-set fixed grid using the 5th order WENO, and the signed distance function is updated the fast marching scheme.

The detailed description of the sub-optimization step other than pre-processing or post-processing operations, are found in the design structure matrix of the sub-optimization shown in Figure 7.

![Design structure matrix of sub-optimization found in LSTO](image)

The workflow of the sub-optimization of the LSTO is composed of 6 Components. The first `input_comp` Component specifies the 4 independent variables: Lagrange multipliers (`lambdas`) as design variables, length segments for boundary integral (`segments`), sensitivities with respect to the objective and constraints (`sensitivities`), and the indices of the active nodes (`active_ids`). Note that not all of nodes in the level-set grid are active during the iteration. For example, the nodes belonging to the domain boundary or non-designable regions and their level-set values remain fixed to their initial value. Filtering the values at active boundary points (`active_comp`) is, therefore, a necessary step in order to reduce the number of degree of freedoms. The subsequent Component computes boundary-integrated values (`integ_comp`) and its corresponding advection distances (`distance_comp`) based on Eq. (8). It is worth pointing out that a finite difference scheme is inevitably used to estimate the partial derivatives of the distances with respect to $\lambda$ because the distances are not linear with respect to the input as the side limit and the CFL condition are possibly violated when the distances are
linearized [7]. Therefore, a numerical differentiation scheme provided by OpenMDAO is utilized in the Component, which also demonstrates the efficiency and flexibility in practical implementations. The Components that follow are dedicated to the objective function (objective_comp) and constraint function (constraint_comp) based on Eq. (9). The resulting advection velocity is expected to be optimal for compliance minimization, while satisfying the constraints.

The resulting advection velocity is expected to be optimal for compliance minimization, while satisfying the constraints.

**IV. Extensions**

In addition to the unique architecture of OpenMDAO, which enhances the modularity of the optimization schemes and the ease of the implementations, using OpenMDAO as a topology optimization platform is also advantageous in extending or modifying the existing implementations. In the present section, we demonstrate the ease of restructuring and the reusability of the modules by example.

**A. Flexible arrangement of the module**

The restructurability of the program is demonstrated by rearranging the order of the density filter Component with respect to other Components. Thanks to the encapsulation of the Component object, minimal changes are required, in

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**Fig. 8** Numerical results of LSTO. (a) a convergence profile with initial configuration where size (Lxy) and the number of meshes (Exy) to each direction are specified (b) a material configuration, area fraction (G) and compliance (c) found during at the 30th, 80th, and 200th iterations.

Figure 8 shows a set of results and the convergence graph for the same cantilevered beam problem solved previously with SIMP. As the hole-creation method [14] is not employed herein, the initial design domain has seeded holes. Both the compliance and constraint values exhibit a smooth convergence as reported in the literature [7], which is in contrast to SIMP example. A topology similar to SIMP is found in the converged optimal solution, although with more holes.

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the Group layer only. This is in contrasts to the topology optimization programs such as the minimalistic educational implementations [9]. No matter how simple the program might be, it is still true that small tweaks such as the removal of the filter require re-programming of the code; not only the functions but also the sensitivities must be changed accordingly.

One of the possible tweaks to the configuration are found in the Figure 9, where the corresponding design structure matrix and code snippet are presented.

![Figure 9](image)

**Fig. 9** A demonstration of restructurability. (a) the design structure matrix of the SIMP method, where the filter object is rearranged with respect to that of the penalization step, and (b) a code snippet for the reconfiguration that exemplifies the easy reconfiguration; commented commands are marked by blue and corresponding changes are marked by red.

When compared with the design structure matrix of SIMP (Figure 5), the Component object for the density filter \((\text{filter}_\text{comp})\) is shown to be rearranged relative to the penalization filter \((\text{penalization}_\text{comp})\) (Figure 9(a)). The code snippet that realizes the rearrangement is also shown in Figure 9(b). The change of the code is proven to be minimal as the reconnection of the inputs and outputs within the Group object is the only required modification, thanks to the \texttt{connect} member function of the Group object and the thorough encapsulation of the Component object. Even though the whole code is not shown herein, one may easily see how the removal of the filter can be realized. It is worth mentioning that these two tweaks are presented only to succinctly demonstrate the possible extensions, and these extensions are not necessarily typical in the SIMP method; however, this example is in line with reducing the repetitive
programming required in selecting the type of weight of the filter.

Fig. 10  Optimal solutions where the filter object is (a) rearranged or (b) removed.

The numerical results are shown in Figure 10. As one may expect, either rearranging (Figure 10(a)) or removal (Figure 10(b)) of the existing filter each saliently generates an intermediate density or a checkerboard pattern in the converged solutions. This example also exhibits the educational benefits, as an effect of the filter as a way to remove the ill-posedness of the problem is demonstrated without the drastic changes of the code.

B. Parameterized level-set topology optimization

The reusability of the modules is also exemplified by extending the present Components to a new topology optimization formulation that bears similarities to the parameterized level-set topology optimization approach [5,22,23]. This method is, in essence, a density-based approach, but the densities are parametrized using a level set function that implicitly represents the boundary. Instead of tracking the boundary directly, the value of this function is used to control the densities in the elements, using a Heaviside function to map to the (0, 1) interval and SIMP-type penalization to discourage intermediate densities. Further investigation is necessary to explore the potential of this method as a new approach for topology optimization, but our goal here is to highlight that the use of OpenMDAO as a topology optimization platform facilitated the development of this approach through reusability and restructurability. As shown in Figure 11, many Components from SIMP are reused, including the implicit Component that computes the displacements using linear elasticity (states_comp), the penalization Component (penalization_comp), and the objective (objective_comp) and constraint function (weight_comp) Components.

In this scheme, design variables are the localized values of the regularly spaced control points, by which a hypersurface is constructed via a parametric mapping using B-splines (parametrization_comp). The generated surface is then filtered into discrete values bound to (0, 1) by passing them into an analytic Heaviside function (heaviside_comp): the hyperbolic tangent function. The filtered values are interpreted as a material density (averaging_comp); if their spatial size coincides to that of a finite element mesh, as in the present case, the SIMP module without a filter can be reused for the remaining computation.

The numerical results are presented in Figure 12. Although the results show oscillations in the boundary, they are sufficient to demonstrate that the optimal topology computed out of the present scheme and its compliance value are comparable to those of SIMP. We note that the reusability demonstrated herein can enhance the productivity in implementation, as new ideas or different modules can be quickly implemented without repeated programming once the
Fig. 11 Design structure matrix of the parametric level-set approach for topology optimization.

Fig. 12 Numerical results computed using the parametric level-set approach for topology optimization: (a) the convergence profile with the initial specifications same as in SIMP (b) the material configuration, area fraction \( G \) and compliance \( c \) at the 4\(^{th} \), 17\(^{th} \), and 90\(^{th} \) iterations.

V. Conclusion

Topology optimization is a structural design method capable of computing the optimal layout among a large number of candidates. There is ongoing research in extending it to multiscale or multidisciplinary applications; however, its potential is not fully exploited due to the non-technical problems associated with the implementation time and effort. This is not only inefficient but also makes the program prone to human-induced errors.

In this work, we incorporate topology optimization into OpenMDAO, an computational framework for MDO. The
primary benefit of using OpenMDAO comes from its modular architecture. By decomposing each topology optimization method into Components, the programming becomes more object-oriented and the derivatives across the multiple Components are automatically calculated. Also, the workflow can be easily visualized as connectivities between Components. Another benefit is easy application of standard nonlinear and linear solvers available in OpenMDAO. We demonstrate these benefits using the two most widely used topology optimization methods, SIMP and the level-set method. We also demonstrate the extensibility by modifying the SIMP implementation to perform parameterized level-set topology optimization, which is a third method. The modular approach enables this modification with minimal additional programming effort where most of the existing Components are reused.

The authors expect the current work to be applied to the different physics as the required change is minimal when compared with using approaches [9]. Moreover, OpenMDAO’s suite of optimization methods and utilities potentially benefits in educating and promoting the topology optimization strategies to researchers in other disciplines.

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