Directional radiometry and radiative transfer: The convoluted path from centuries-old phenomenology to physical optics

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ABSTRACT

This Essay traces the centuries-long history of the phenomenological disciplines of directional radiometry and radiative transfer in turbid media, discusses their fundamental weaknesses, and outlines the convoluted process of their conversion into legitimate branches of physical optics.

1. Introduction

It is a great honor to have become the second recipient of the Hendrik C. Van de Hulst Award presented by Elsevier in the general category of Electromagnetic Scattering. It is also a special pleasure for me to receive this award following Prof. Joop W. Hovenier (Fig. 1) who has always exerted strong influence on my research and had unknowingly served as an implicit adviser during my PhD studies and early years in science. Part of the award ceremony was a Van de Hulst Lecture presented at the 14th Conference on Electromagnetic and Light Scattering (ELS) on 20 June 2013. Another official part of this award is the honor and obligation to publish in the Journal of Quantitative Spectroscopy and Radiative Transfer (JQSRT) a scientific Essay intended to summarize the recipient's personal view of the state-of-the-art of one or more disciplines related to electromagnetic scattering by particles and particulate media. Needless to say, these disciplines are expected to have been foci of the recipient's own research leading to the Van de Hulst Award.

The formats and styles of the Van de Hulst Lecture and the Van de Hulst Essay are still in a state of flux since only two such lectures have been presented and only one such Essay has been published [1]. On one hand, this lack of established traditions appears to represent a challenge. Yet on the other hand I feel that it sets me free to adopt an ad hoc format for this Essay and essentially speak my mind with the hope that the result will be instructive to the JQSRT readership. Therefore, I have decided to focus on the current state of the disciplines of directional radiometry and radiative transfer as I understand them following...
The 1998 ELS conference was convened at the NASA Goddard Institute for Space Studies (GISS) in New York, and Professor Van de Hulst had kindly agreed to open it with a keynote lecture. He was very enthusiastic about the occasion to visit the place where he spent a six-month sabbatical in 1962, the result of which was his famous NASA report on the adding/doubling method [5]. Unfortunately, just two days before the opening of the conference Professor Van de Hulst faxed me with an apology for being unable to come to New York because of certain health issues. However, he was still able to contribute the instructive Foreword [6] to the monograph on light scattering by nonspherical particles [7] which has proved to be an important collective outcome of the New York conference.

Since Joop Hovenier did his PhD work under the supervision of Professor Van de Hulst and since my early research had benefitted so much from Joop's publications (especially the 1983 review [8] co-authored by Cornelis van der Mee), I consider myself one of Van de Hulst's "indirect" disciples. My direct scientific genealogy can be traced to Academician Viktor Ambartsumian, one of the founders of theoretical astrophysics. Indeed, I did my PhD work under the supervision of Dr. Edgard Yanovitskij whose PhD thesis was in turn supervised by Academician Sobolev (Fig. 4), the prominent Soviet astrophysicist and the best known PhD student of Academician Ambartsumian (Fig. 5). My early research was also influenced by two other members of the Ambartsumian–Sobolev school of radiative transfer, Vsevolod Ivanov and Helmut Domke (Fig. 6). Furthermore, Vsevolod Ivanov served as an Official Opponent on my PhD and Habilitation thesis defenses.

As already mentioned, I owe my initial basic knowledge of the theory of radiative transfer to Van de Hulst's monograph [3], as well as to the so-called “blue Sobolev” [9]. That nickname refers to the navy blue color of the cover in the original Russian edition of this well-known monograph and was used by Soviet scientists to distinguish it casually from the “black Sobolev” [11]. As a novice in the field of radiative transfer, I took for granted the apparent simplicity and obviousness of the main phenomenological concepts of this discipline and for some time had not realized that the "traditional" radiative transfer theory (RTT) is, figuratively speaking, a "colossus with feet of clay". My awakening started in 1986 when I read the Russian edition of the 1978 book by Ishimaru [12]. It was quite surprising to learn that

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*Fig. 1. Joop Hovenier (left) and Michael Mishchenko at ELS-XIV in Lille on 17 June 2013.*
there appeared to be at least two different and completely unrelated ways of arriving at exactly the same scalar RTE for a turbid medium, **viz.**, the "traditional" back-of-an-envelope derivation allegedly based on the concept of radiance and energy conservation considerations as well as the one allegedly based on the scalar wave theory. As someone with a bit of a background in mathematical logic, I thought that if that were the case then one of the derivations must be a direct corollary of the other because otherwise it would be a fake derivation essentially amounting to the postulation of the RTE based on verbal "simple physical considerations".

The following 25 years of study and research have confirmed that initial suspicion. Furthermore, I believe that this research has served to conclude the work initiated by Rudolph Preisendorfer, Yuri Barabanenkov, Anatoli Borovoi, Yuri Gnedin, Emil Wolf, Akira Ishimaru, Leung Tsang and others in that it has brought the disciplines of directional radiometry and radiative transfer into the realm of physical optics. Therefore, what follows is a personal and, by definition, subjective account of what it has taken to finally convert the centuries-old phenomenologies of directional radiometry and radiative transfer, as applied to particulate media, into first-principle theories.

### 2. Phenomenology

A thorough account of the early history of directional photometry\(^2\) was provided by DiLaura in the introduction to Ref. [13]. He attributed the culmination of medieval optics to

\(^2\) For the purposes of this Essay, the terms "photometry" and "radiometry" will be used interchangeably.
Ad Vitellionem Paralipomena [14] by Johannes Kepler (1571–1630; Fig. 7) published in 1604 and containing one of the most fundamental elements of photometry, viz., the attenuation of the intensity of light as the inverse square of distance from a point-like source. However, establishing photometry as a scientific discipline was the outcome of systematic studies by the French natural scientist and engineer Pierre Bouguer (1698–1758) followed by those of the Swiss mathematician, natural scientist, and philosopher Johann Lambert (1728–1777).

Bouguer′s Essai d′Optique [15] was published in 1729, while its thorough augmentation, Traité d′Optique [16] (Fig. 8), appeared posthumously in 1760. Bouguer′s research was mostly experimental and relied on several ingeniously designed photometric instruments. He was the first to realize that the human eye cannot be used as an accurate absolute meter of brightness but is quite capable of establishing the equality of brightness of two adjacent surfaces. In Essai d′Optique Bouguer describes the use of Kepler′s law of inverse squares and the human eye as an equality indicator to derive the ratio of luminous intensities of two light sources and discovers the famous exponential attenuation law\(^3\) by studying the diminution of light as it passes through translucent media (Fig. 9). He also analyzed the reflection of light by rough surfaces and proposed the idea of modeling such diffuse reflectors as consisting of small randomly oriented mirrors whose orientation distribution would determine the macroscopic angular reflectance. In his Photometria [17] (Fig. 8), Lambert was the first to extensively and systematically use contemporary mathematics, including calculus, to interpret experimental results and developed the mathematical foundation of radiometry by introducing specific definitions of photometric quantities and a unified set of photometric principles and laws.

The impact of Bouguer′s and Lambert′s work was so profound that even now much of illumination engineering is based, directly or indirectly, on their treatises. Perhaps the only significant augmentation dating from 1854 was the incorporation of the solution concentration into Bouguer′s exponential attenuation law by August Beer (1825–1863) [18].

With the development of Maxwell′s electromagnetics in 1864 and the realization that light consists of electromagnetic waves, the photometry of Bouguer and Lambert

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\(^3\) Bouguer′s exponential attenuation law is often incorrectly attributed to Lambert. This is thoroughly inappropriate since Lambert had read Bouguer′s Essai d′Optique published in 1729 and frequently cited it in his own Photometria [17] published in 1760.
could no longer be considered a branch of physics based on first principles and had thus become a purely phenomenological discipline. However, this profound and irreversible shift in the conceptual status of the old directional photometry has largely been ignored. All in all, phenomenological photometry, as summarized eloquently in 1936 by Andrei Gershun (1903–1952) in terms of the so-called “light field” [21], has been one of the oldest surviving paradigms in contemporary science despite its whopping disconnect from the “mainland” of modern physics (in the words of Rudolph Preisendorfer [28]).

The main contribution by the German physicist and mathematician Eugen von Lommel (1837–1899) was to introduce, in 1887, the notion of the amount of radiant energy crossing an imaginary geometrical rather than an actual physical surface element. This allowed him to conceptualize the directional flow of radiant energy through space and introduce the integral form of the RTE as a way of solving the problem of diffusion of light through a turbid medium composed of isotropically scattering centers [29] (Fig. 10). Virtually identical results were published independently by the Russian physicist Orest Khvolson (1852–1934) two years later [30] (Fig. 10; see also the instructive account of the early history of the phenomenological RTT by Ivanov [31]).

The work by Lommel and Khvolson has remained largely unnoticed. The first introduction of the RTE has traditionally been attributed to Arthur Schuster (1851–1934). In actuality, however, Schuster’s paper of 1905 [32] contains what is now known as the two-stream approximation rather than the integral or integro-differential form of the RTE. The first phenomenological derivation of the RTE in the case of anisotropic scattering was given by Louis Vessot King in 1913 [33]. Like Lommel and Khvolson, King introduced the integral form of the RTE rather than the integro-differential equation that eventually assumed the status of being the canonical form of the RTE. Interestingly, the direct microphysical derivation from the Maxwell equations yields the integral form of the RTE, the integro-differential form being a corollary.

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4 A physical theory is called phenomenological if it expresses mathematically the results of observed phenomena without clarifying their fundamental origin and significance. Typically, the development of a phenomenological theory is based on experience-based heuristic shortcuts lacking rigorous justification. Most phenomenological theories are short-lived and get replaced by fundamental first-principle theories. However, as we discuss in this Essay, some phenomenologies can survive for centuries despite their inherently limited scientific value and eventually become an impediment to scientific progress. The Latin term “Phenomenologia” was introduced by the German Lutheran theologian and theosopher Christoph Friedrich Dettlinger in 1736 [19]. Subsequently, the German term “Phänomenologie” was used by Lambert to name his “doctrine of appearance” [20]. It should be noted that Lambert viewed his photometry as a phenomenological discipline from the very outset.

5 The concepts of scientific paradigms and paradigm shifts emerged in the framework of the historical approach in the philosophy of science according to which only by studying history of science can we gain an adequate understanding of human reason. These profound notions were introduced and analyzed, using different terminology, by the 20th century French philosophers Gaston Bachelard, Georges Canguilhem, and Michel Foucault, but the roots of their work can be traced to the older French tradition of science studies, going back to Auguste Comte and including later thinkers such as Pierre Duhem and Henri Poincaré (see, e.g., Refs. [22–26]). Thomas Kuhn [27] systematized and popularized these ideas and re-instituted the use of the term “paradigm” introduced by Plato in his dialog Timaeus to name the eternal pattern used by a divine Craftsman (the Demiurge) to create the universe. A universally accepted definition of a scientific paradigm hardly exists, but in general it can be summarized as a self-contained set of concepts, values, perceptions, and practices shared by a large scientific community; it forms a particular vision of reality and is the basis of the way the community organizes itself. A paradigm shift (or, in Kuhn’s terminology, a scientific revolution) occurs when a given discipline switches from one distinct paradigm to another, often as a result of a long process.

6 The first page of Ref. [30] contains an interesting footnote by Academican H. Wild according to which the paper was originally submitted in the fall of 1885. It was then withdrawn by the author who hoped to obtain a more complete solution of the main equation. The paper was resubmitted in the fall of 1888 essentially in its original form. It thus appears that Lommel and Khvolson introduced the RTE independently of each other.
The key quantity of the phenomenological photometry and phenomenological RTT is the specific intensity (also called radiance), which is postulated to have primordial physical existence and a priori defined properties. The standard definition of the specific intensity was given in 1906 by Max Planck (1858–1947) in his famous *Theorie*...
The state of the radiation at a given instant and at a given point of the medium cannot be represented by a single vector (that is, a single directed quantity). All heat rays which at a given instant pass through the same point of the medium are perfectly independent of one another, and in order to specify completely the state of the radiation the intensity of radiation must be known in all the directions, infinite in number, which pass through the point in question.

Based on this premise, the monochromatic radiance $I(r, \hat{q})$ is defined by stating that the amount of monochromatic radiant energy $dE$ transported through an arbitrarily chosen differential element of area $dS$ in the interior of a medium in directions confined to a differential element of solid angle $d\Omega_{\hat{q}}$, centered around the propagation direction $\hat{q}$, during a differential time interval $dt$ is given by

$$dE = I(r, \hat{q}) \cos \theta \, dS \, dt \, d\Omega_{\hat{q}}, \quad (1)$$

where $r$ is the position vector of the differential surface element and $\theta$ is the angle between $\hat{q}$ and the normal $\hat{n}$ to $dS$ (Fig. 12a). This definition was eventually adopted in the classical works by E. Arthur Milne [36], Eberhard Hopf [37], and Subramanyan Chandrasekhar [38] as well as in virtually all subsequent monographs and textbooks on the RTT and directional radiometry (see, e.g., Refs. [9,11,39–61]). Although in his treatise Planck specifically considered black-body electromagnetic radiation, his concept of the specific intensity was extended to encompass the scattering of light by cloudy and
other particulate media, which is the specific subject of this Essay. The heuristic notion of the radiance was eventually supplemented by the traditional belief that it can be directly measured with a suitable optical device such as the so-called Gershun tube sketched in Fig. 13 [21,47,55].

The French physicist and astronomer (as well as the 25th Prime Minister of France) Dominique François Arago (1786–1853) was among the first to criticize phenomenological radiometry for complete ignorance of polarization state of light. This criticism was nominally addressed by replacing the radiance $I(r, q)$ with the four-element specific intensity column vector $\mathbf{I}(r, q)$. In 1924, Richard Gans (1880–1954) considered the transfer of polarized light in a plane-parallel Rayleigh-scattering atmosphere [62], but analyzed only the special case of perpendicularly incident light and considered only the first two components of the specific intensity column vector. The case of arbitrary illumination and arbitrary polarization was addressed in 1950 by Subrahmanyan Chandrasekhar (1910–1995) [38].

In 1955, Georgi Rozenberg (1914–1982) introduced the most general form of the integro-differential vector RTE applicable to sparse scattering media composed of arbitrarily shaped and arbitrarily oriented particles [63]. This publication (see also Ref. [64]) had essentially concluded the conceptual development of the phenomenological RTT. Perhaps the only significant subsequent augmentation was the inclusion of the thermal emission vector in the Rozenberg’s vector RTE by Leung Tsang in 1984 [65] (corrected in Ref. [66]).

The work by E. Arthur Milne (1896–1950), Eberhard Hopf (1902–1983), Viktor Ambartsumian (1908–1996), Subrahmanyan Chandrasekhar, Viktor Sobolev (1915–1999), and Hendrik van de Hulst (1918–2000), among others, had served to establish the phenomenological RTT as a branch of mathematical physics (see Refs. [67–75] and references therein). Most recently, the phenomenological RTT has been incorporated into the equally phenomenological discipline of computer graphics studying techniques to digitally synthesize and manipulate images (see, e.g., Refs. [76–78] and references therein).

3. What is fundamentally wrong with the phenomenological approach to directional radiometry and RTT?

3.1. Polydirectional flow of radiant energy

In the introduction to the English translation of Gershun’s treatise [21], Moon and Timoshenko wrote in 1939:

Theoretical photometry constitutes a case of “arrested development”, and has remained basically unchanged since 1760 while the rest of physics has swept triumphantly ahead. In recent years, however, the increasing needs of modern lighting technique have made the absurdly antiquated concepts of traditional photometric theory more and more untenable.

The Gershun’s treatise was then presented as part of “a vigorous attempt to bring the theory of light calculation into conformity with the spirit of physics.” However, some 70 years after Maxwell’s Dynamical Theory [79], Gershun’s way of conforming with Maxwell’s electromagnetics was, in fact, to ignore it. Instead, the centerpiece of his treatise is the so-called “light field”, i.e., “a part of space studied from the standpoint of transmission of radiant energy.
within that space.” To justify this approach, Gershun wrote in the introduction to his book:

A reader-physicist would naturally ask why the author distinguishes the light field from the electromagnetic field well studied in physics. It is true that the light field is caused by the electromagnetic field, but qualitatively it is quite different.7

After this programmatic statement, electromagnetics is hardly mentioned in the rest of Ref. [21]. This manifesto, according to which the radiance field is somehow qualitatively different from the electromagnetic field and thus must be studied using different physical principles, has been implicit in the majority of “phenomenological” publications on directional radiometry and radiative transfer.

Whether spelled out explicitly or not, the key premise of phenomenological photometry as well as of the phenomenological RTT is that matter interacts with the energy of the electromagnetic field rather than with the electromagnetic field itself. This profoundly false assumption explains the deceitful simplicity of the phenomenological electronics as well as their ultimate failure. Indeed, the very outset of both phenomenological disciplines is the postulation of the existence of the radiance as the primordial physical quantity describing the “instantaneous directional distribution of the radiant energy flow” at a point in space. This is followed by a “derivation” of the scalar RTE on the basis of “simple energy conservation considerations” and the postulation that it is the electromagnetic energy rather than the electromagnetic field that gets scattered by particles and surfaces.

However, it is imperative to recognize that according to classical electromagnetics, the field–matter interaction is controlled by the electric and magnetic field vectors rather than by the energy of the electromagnetic field [80,81]. Similarly, the canonical formulation of classical electrodynamics is based on expressing the field–matter interaction term in the Lagrangian and Hamiltonian densities in terms of the electromagnetic 4-vector potential, which, in turn, is again related to the electric and magnetic fields. The canonical formulation of quantum electrodynamics (QED) is obtained by promoting the electromagnetic 4-vector potential and the current density of matter to field operators that satisfy specific commutation relations [82–88]. This implies that any attempt to bypass the explicit solution of the Maxwell equations or an explicit QED computation in a first-principle development of the RTT and thus build the RTT on the notion of radiant energy rather than on the notion of the electromagnetic field (see, e.g., Ref. [89]) is fundamentally flawed and is doomed from the very outset.

Furthermore, the very notion of polydirectional propagation of electromagnetic energy at a point in space, as allegedly described by the radiance, contradicts basic laws of classical electromagnetics and does not follow from QED. Indeed, the quantity characterizing instantaneous electromagnetic energy transport in classical electromagnetics is the Poynting vector \( \mathbf{S} \) given by the vector product of the real-valued electric, \( \mathbf{E} \), and magnetic, \( \mathbf{H} \), field vectors:

\[
\mathbf{S}(\mathbf{r}, t) = \mathbf{E}(\mathbf{r}, t) \times \mathbf{H}(\mathbf{r}, t),
\]

where \( \mathbf{r} \) is the position vector and \( t \) is time [80,81]. However, by virtue of being a direct corollary of the Maxwell equations, the famous Poynting theorem involves the integral of the Poynting vector over a closed surface. As such, it quantifies the energy budget of a finite volume element rather than the local flow of electromagnetic energy at a point in space. Back in 1916, the Dutch physicist Hendrik Antoon Lorentz (1853–1928) already warned against too literal an interpretation of the Poynting vector as describing a current of electromagnetic energy by noting that

in general it will not be possible to trace the paths of parts or elements of energy in the same sense in which we can follow in their course the ultimate particles of which matter is made up. …It might even be questioned whether, in electromagnetic phenomena, the transfer of energy really takes place in the way indicated by Poynting’s law (see pages 25–26 of The Theory of Electrons [90]). Even if the Poynting vector could be claimed to characterize the local current of electromagnetic energy, this vector is inherently monodirectional at any moment in time and remains monodirectional upon averaging over any time interval. Therefore, there is no reason whatsoever to

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7 This translation is somewhat different from that by Moon and Timoshenko and, in my opinion, is more accurate.
postulate that there exists a primordial physical quantity, viz., the radiance, that characterizes the polydirectional local flow of electromagnetic energy.

3.2. Photonic confusion

The often uncontrollable use of the word "photon" taken out of its proper QED framework is another manifestation of the implicit desire to bypass the complexities of dealing with the electromagnetic field. Indeed, a popular way to justify the primordial existence of the polydirectional radiance is to claim that the light field is in fact a "gas" of localized point-like particles of light called photons. Usually this is done with reference to Albert Einstein (1879–1955) who attempted to model the photoelectric effect heuristically by resurrecting the idea of light corpuscles advocated by Sir Isaac Newton (1643–1727). Specifically, Einstein suggested in 1905 that

the energy of a light ray spreading out from a point source is not continuously distributed over an increasing space but consists of a finite number of energy quanta which are localized at points in space, which move without dividing, and which can only be produced and absorbed as complete units.
(see Ref. [91]). Since such localized point-like photons must exist in vast numbers, one can imagine that at any given moment many photons can fly by a point in space in different directions, thereby representing the sought polydirectional localized transport of radiant energy [92–96].

The photons are not allowed to collide with each other, but they are allowed to collide with macroscopic particles such as cloud droplets. In the words of Kirk [46], “the photons follow a zig-zag path as they ricochet from one scattering particle to the next.” These “ricochets” are random and are claimed (again, on the basis of “simple physical considerations”) to lead to a Boltzmann type kinetic equation, viz., the scalar RTE. Unfortunately, it has never been explained why the kernel of this equation, which by design describes the “ricochet” of a photon from a macroscopic particle, is always calculated by solving the classical macroscopic Maxwell equations (MMEs) which involve neither photons nor “photon–particle collisions.” Apparently, it is believed that upon approaching a droplet the photon becomes an electromagnetic wave. This allows the photon to “ricochet” from the droplet according to the MMEs (e.g., according to the formulas of the Lorentz–Mie theory) and become an outgoing spherical wave. Eventually, the outgoing spherical wave switches back to being a bunch of localized point-like photons. Needless to say, such mysterious transformations of photons into waves and then back into photons can only happen in the confused human mind rather than in nature. The only outcome of such verbal speculations can be – and has been – the notorious “photonic confusion”.

The lasting misinterpretation of the actual QED photons as localized particles of light has been kept flourishing by scores of incompetent popularizers of science, contributors of amateur Wikipedia articles, and authors of many school and college textbooks and even professional monographs. A typical example is the textbook by Taylor et al. [97]. On p. 139, one can find the following misleading statement: “Today all physicists accept that the photoelectric effect, the Compton effect, and numerous other experiments demonstrate beyond doubt the particle nature of light.” This statement ignores the well-established fact that the alleged particle behavior of light in phenomena such as the photoelectric effect can be explained quantitatively in terms of the semi-classical approach wherein the electromagnetic field is not quantized and is described by the classical microscopic Maxwell equations [98–100]. Furthermore, based on the advanced QED theory of the photoelectric effect, Kimble and Mandel [101] concluded that photodetectors do not count photons in any precise sense. For some reason these facts are virtually never mentioned in school and college textbooks. Another profoundly wrong state-

However, the obsolete heuristic nature of Einstein’s localized light quanta [104–106] becomes patently obvious upon opening an advanced textbook on the QED or quantum optics (e.g., Refs. [107–110]). Indeed, although the term “photon” is ubiquitous in those disciplines, it is well recognized that there is no position operator for an actual QED photon and that it is impossible to define a photon wave function in the coordinate representation, which precludes photon localizability in space (e.g., Section 2.2 of Ref. [83]). The real QED photons are quantum excitations of the normal modes of the electromagnetic field and as such are associated with electromagnetic plane waves of definite wave vector and definite polarization but infinite lateral extent. These factors imply that QED photons are not localized point-like particles of light and as such cannot be used to justify the notion of polydirectional local flow of radiant energy allegedly described by the specific intensity [111].

Some 85 years since the development of QED by Paul Adrien Maurice Dirac, Ernst Pascual Jordan, and Werner Karl Heisenberg [112–115], the superficial use of the word “photon” is still frequently accompanied by the mention of the alleged “wave – particle duality” of light. It is therefore instructive to quote from the famous 1995 Anti-Photon by Willis Lamb Jr. [106], where he notes that talking about the wave – particle duality in discussion of quantum mechanics ... may be necessary for those who are unwilling or unable to acquire an understanding of the theory. However, this concept is even more pointless introduced in discussions of problems in the quantum theory of radiation.

He concludes the Anti-Photon by stating that

It is high time to give up the use of the word “photon”, and of a bad concept which will shortly be a century old. Radiation does not consist of particles, and the classical, i.e., non-quantum, limit of the quantum theory of radiation is described by Maxwell’s equations for the electromagnetic field, which do not involve particles. Talking about radiation in terms of particles is like using such ubiquitous phrases as “You know” or “I mean” which are very much to be heard in some cultures.9

In 1989, Kidd et al. [105] suggested that elementary texts would benefit from dropping the corpuscular photon (except, perhaps, as a historical topic) and switching to the semi-classical treatment as the first approximation to the modern QED approach. In 1995, Lamb Jr. [106] wrote that “the sooner an appropriate reformulation of our educational processes can be made, the better.” However, it

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9 I must admit my own “guilt” of misusing the word “photon” in several early publications. A typical example is the physically meaningless expression “weak localization of photons” used in lieu of the correct (but less catchy) expression “weak localization of electromagnetic waves.”
appears that imaginary point-like photons continue to proliferate, while the inability to explain the nature of the actual QED photon is still disguised by references to the mysterious “wave–particle duality” of light.

3.3. Preisendorfer’s radiance function

The need to establish a conceptual link between the “island” of the phenomenological RTT and the “mainland of electromagnetic theory” was eloquently formulated in 1965 by Rudolph Preisendorfer (see Chapter XIV of Ref. [28]). He admitted that the heuristic polydirectional radiance cannot be defined as existing at a specific moment \( t \) but rather must be the result of averaging over a sufficiently long time interval. He observed that in a turbid medium, the constituent particles are in constant motion and can also change their sizes, shapes, and orientations, thereby rendering the direction and magnitude of the Poynting vector \( \mathbf{S}(\mathbf{r}, t) \) at an observation point \( \mathbf{r} \) random functions of time. At certain moments the direction of the instantaneous Poynting vector can fall within the differential solid angle \( d\Omega_{q} \) in Fig. 12a. Therefore, Preisendorfer suggested that an appropriate definition of the specific intensity could be the time-averaged length of the Poynting vectors at \( \mathbf{r} \) with directions falling within \( d\Omega_{q} \). More specifically, Preisendorfer’s radiance function is defined as follows (see Fig. 12a):

\[
\bar{N}(\mathbf{r}, \hat{q}) = -\frac{1}{\cos \theta} \lim_{\Delta S \to 0} \lim_{\Delta \Omega_{q} \to 0} \lim_{t \to \infty} \frac{1}{\Delta S} \int_{\Delta S} d^{2}r \times \int_{t-(T/2)}^{t+(T/2)} d\tau \langle |\mathbf{S}(\mathbf{r}, \tau)|/\mathbf{S}(\mathbf{r}, \tau) \rangle [\chi[\Delta \Omega_{q}, \hat{S}(\mathbf{r}, \tau)]],
\]

where \( \hat{S}(\mathbf{r}, \tau) = \mathbf{S}(\mathbf{r}, \tau)/|\mathbf{S}(\mathbf{r}, \tau)| \) is the unit vector in the direction of \( \mathbf{S}(\mathbf{r}, \tau) \) and

\[
\chi[\Delta \Omega_{q}, \hat{S}(\mathbf{r}, \tau)] = \begin{cases} 
1 & \text{if } \hat{S}(\mathbf{r}, \tau) \in \Delta \Omega_{q} \\
0 & \text{otherwise}
\end{cases}
\]

is the angular step function. Essentially the same definition of the radiance can be found in Section 7.9 of Ref. [12]. Preisendorfer then attempted to use the Maxwell equations to demonstrate that his radiance function \( \bar{N}(\mathbf{r}, \hat{q}) \) satisfies the scalar RTE.

However, Preisendorfer’s attempt to bridge the gap between the phenomenological RTT and Maxwell’s electromagnetics failed for several reasons. First of all, we have already mentioned that there is no fundamental reason to believe that the Poynting vector specifies the direction and magnitude of the instantaneous local flow of electromagnetic energy. Secondly, the derivation of the RTE in Ref. [28] turned out to be incorrect owing to the wrong underlying assumption that the instantaneous electric and magnetic field vectors at any point inside a turbid medium are always mutually orthogonal. Thirdly, it can be demonstrated that the radiance function \( \bar{N}(\mathbf{r}, \hat{q}) \) does not satisfy the RTE [116]. Fourthly, even if \( \bar{N}(\mathbf{r}, \hat{q}) \) were to satisfy the RTE, this quantity would be useless because it cannot be measured at a point inside a volume of turbid medium.

To illustrate the last statement, let us assume that there exists a hypothetical instrument that reacts to the magnitude of the instantaneous local Poynting vector only if the direction of \( \mathbf{S}(\mathbf{r}, t) \) falls within a narrow acceptance solid angle \( \Delta \Omega_{q} \) where \( \mathbf{r} \) is a point on the sensitive surface \( S \) and the unit vector \( \hat{q} \) specifies the orientation of the optical axis of the instrument (Fig. 12b). Let us also assume that this hypothetical instrument is placed inside a random cloud consisting of \( N \) particles (Fig. 14). It is quite obvious that averaging the reading of such an instrument over a sufficiently long period of time would essentially yield the Preisendorfer’s radiance function \( \bar{N}(\mathbf{r}, \hat{q}) \) provided that the presence of the instrument does not affect the instantaneous Poynting vector \( \mathbf{S}(\mathbf{r}, t) \) for any \( \mathbf{r} \in S \), where \( \mathbf{r} \) is the position vector of the central point of the sensitive surface.

However, it is easily seen that no matter how small the instrument is relative to the cloud, its very presence serves to not just affect, but completely destroy the quantity that is supposed to react to [117]. Indeed, let us assume for simplicity that the cloud particles are separated widely enough to satisfy the conditions of applicability of the far-field Foldy equations (Section 4.5). Then the total instantaneous electric and magnetic fields at \( \mathbf{r} \) in the absence of the detector are superpositions of the respective incident and \( N \) partial scattered fields:

\[
\mathbf{E}(\mathbf{r}, t) = \mathbf{E}^{inc}(\mathbf{r}, t) + \sum_{i=1}^{N} \mathbf{E}^{sc}(\mathbf{r}, t),
\]

\[
\mathbf{H}(\mathbf{r}, t) = \mathbf{H}^{inc}(\mathbf{r}, t) + \sum_{i=1}^{N} \mathbf{H}^{sc}(\mathbf{r}, t),
\]

where \( \mathbf{E}^{sc}(\mathbf{r}, t) \) and \( \mathbf{H}^{sc}(\mathbf{r}, t) \) describe an outgoing spherical wavelet centered at the origin of particle \( i \). By definition, the corresponding local instantaneous Poynting vector is given by the vector product \( \mathbf{S}(\mathbf{r}, t) = \mathbf{E}(\mathbf{r}, t) \times \mathbf{H}(\mathbf{r}, t) \). The major side effect of the presence of the hypothetical detector is to block the spherical wavelets generated by the \( N \) particles located to the left of the plane through the sensitive surface shown schematically by the dashed line in Fig. 14. The resulting “truncated” electric and magnetic fields at \( \mathbf{r} \) are now given by

\[
\mathbf{E}(\mathbf{r}, t) = \sum_{i=1}^{N-N} \mathbf{E}^{sc,i}(\mathbf{r}, t),
\]

\[
\mathbf{H}(\mathbf{r}, t) = \sum_{i=1}^{N-N} \mathbf{H}^{sc,i}(\mathbf{r}, t),
\]

where the sums include only the contributions from the \( N-N \) “unblocked” particles, and we assume for simplicity that the instrument blocks the incident plane wave as well. It is patently obvious that the corresponding “truncated” Poynting vector is not the same as the original Poynting vector:

\[
\mathbf{S}(\mathbf{r}, t) = \mathbf{E}(\mathbf{r}, t) \times \mathbf{H}(\mathbf{r}, t) \neq \mathbf{S}(\mathbf{r}, t).
\]

Irrespective of its potential practical utility, the directional detector of electromagnetic energy flow sketched in Fig. 12b has never been built, and it remains unknown whether it can be designed in principle. Obviously, appraising the very feasibility of building a detector with directional sensitivity to the local instantaneous Poynting vector requires an advanced QED analysis of light–matter interactions.
interaction. It is not inconceivable that there exists a Heisenberg uncertainty relation fundamentally prohibiting such a measurement.

3.4. Directional radiometers

Another profound misconception of phenomenological radiometry is the belief that instruments like the Gershun tube (Fig. 13) measure the directional flow of electromagnetic energy, i.e., filter out the energy propagating within the acceptance solid angle

\[
\Delta\Omega = \frac{\pi D_{en}^2}{4L^2},
\]

where \(D_{en}\) is the diameter of the entrance aperture and \(L\) is the length of the tube (the diameter of the exit aperture \(D_{ex}\) is assumed to be much smaller than \(D_{en}\)). Instead, an elementary physical-optics analysis shows that the Gershun tube filters out plane or near plane electromagnetic wavefronts to propagate within the corresponding instantaneous electric vectors \(E_1\) and \(E_2\) oscillating perpendicularly to the paper, and fully coherent in that at any moment in time \(E_1 = E_2\) at the central point of the entrance aperture. Let the local instantaneous magnetic vectors of the waves be \(H_1\) and \(H_2\), respectively, as shown by the magenta arrows, while the corresponding instantaneous electric vectors \(E_1 = E_2\) are directed towards the reader. The cumulative local instantaneous field is given by the vectors \(E = 2E_1\) and \(H = H_1 + H_2\), the former again being directed towards the reader. One can see that the resulting local instantaneous Poynting vector \(S = E \times H\), shown by the green arrow, is directed along the optical axis of the Gershun tube. Moreover, it is easily verified that the Poynting vector at the central point is always directed along the optical axis of the instrument. Yet the reading of the end photodetector is identically equal to zero since neither truncated wavefront can reach the exit aperture.

This analysis implies that the Gershun tube does not necessarily react to the local Poynting vector at a point in the entrance aperture even if this vector is directed along the optical axis of the instrument. To demonstrate this, let us consider the electromagnetic field formed by a superposition of two plane electromagnetic waves propagating in directions \(\hat{q}_1\) and \(\hat{q}_2\) such that both form a 45° angle with the optical axis of the instrument (Fig. 12d). The waves are linearly polarized, with their electric vectors \(E_1\) and \(E_2\) oscillating perpendicularly to the paper, and fully coherent in that at any moment in time \(E_1 = E_2\) at the central point of the entrance aperture. Let the local instantaneous magnetic vectors of the waves be \(H_1\) and \(H_2\), respectively, as shown by the magenta arrows, while the corresponding instantaneous electric vectors \(E_1 = E_2\) are directed towards the reader. The cumulative local instantaneous field is given by the vectors \(E = 2E_1\) and \(H = H_1 + H_2\), the former again being directed towards the reader. One can see that the resulting local instantaneous Poynting vector \(S = E \times H\), shown by the green arrow, is directed along the optical axis of the Gershun tube. Moreover, it is easily verified that the Poynting vector at the central point is always directed along the optical axis of the instrument. Yet the reading of the end photodetector is identically equal to zero since neither truncated wavefront can reach the exit aperture.

Thus, by its very design, the Gershun tube is a wavefront filter with an acceptance solid angle given by Eq. (10). If this angle is sufficiently small then the Gershun tube can be said to be a well-collimated radiometer (WCR) in that it filters out only those plane or near-plane wavefronts that propagate in essentially the same direction given by the optical axis of the instrument.

In fact, the Gershun tube is a very inefficient WCR since its energy collection efficiency is defined only by the area of the exit aperture \(\pi D_{ex}^2/4\). This area can be increased by increasing \(D_{ex}\), but then the acceptance solid angle \(\Delta\Omega\) also increases and becomes partially vignette. Most WCRs in use today are based on a different optical design, as illustrated schematically in Fig. 12c. Now the main functional elements are the objective and relay lenses, the diaphragm, and the end photodetector. Let us consider the response of such a lens-based instrument to the electromagnetic field formed by superposing two plane waves propagating in directions \(\hat{q}_1\) and \(\hat{q}_2\), respectively. The objective lens acts as a linear optical transformer in that its effect on the total field is a superposition of its effects on each plane-wave component. Specifically, the well-known paraxial approximation (see, e.g., Section 5.1 of Ref. [119]) implies that in the near zone of the objective lens either plane wavefront is transformed into a converging spherical wavefront with its respective focal point located in the plane of the diaphragm. However, the ultimate fates of the two spherical wavefronts in Fig. 12c are different. The pink spherical wavefront passes freely through the pinhole, is converted back into a plane wavefront, and is relayed onto the sensitive surface of the photodetector, thereby contributing to the cumulative reading of the WCR. The blue spherical wavefront gets annihilated by the diaphragm and does not contribute to the photoelectric signal.
Thus the combination [objective lens, diaphragm] serves to select only plane (or near-plane) wavefronts propagating in directions very close to the optical axis of the instrument and falling within its small acceptance solid angle
\[
\Delta \Omega = \frac{d^2}{4f}, \tag{11}
\]
where \(d\) is the diameter of the pinhole and \(f\) is the focal length of the objective lens. The key advantage of this design is that the energy-collection efficiency, defined by the diameter of the objective lens, and the acceptance solid angle are now independent of each other, so that the former can be increased without degrading the latter.

Thus, contrary to a widespread belief, a WCR cannot be said in general to measure the directional distribution of the electromagnetic energy flow at an observation point. It is therefore imperative to formulate precisely what a WCR does in actuality. Let us assume that the lens-based WCR shown in Fig. 12e is exposed to an electromagnetic field in the form of a superposition of several plane wavefronts, as depicted schematically in Fig. 12f. According to the above discussion, the instrument does the following:

- selects only the wavefronts with propagation directions falling within its small acceptance solid angle \(\Delta \Omega \) (i.e., the wavefronts propagating in the directions \(\mathbf{q}_3\), \(\mathbf{q}_4\), and \(\mathbf{q}_5\), but not in the directions \(\mathbf{q}_1\), \(\mathbf{q}_2\), \(\mathbf{q}_6\), and \(\mathbf{q}_7\));
- sums up the respective instantaneous electric and magnetic field vectors: \(\mathbf{E} = \mathbf{E}_3 + \mathbf{E}_4 + \mathbf{E}_5\) and \(\mathbf{H} = \mathbf{H}_3 + \mathbf{H}_4 + \mathbf{H}_5\); and finally
- integrates the modulus of the vector product \(\mathbf{E} \times \mathbf{H}\) (which, by its very construct, is always directed along – or very close to – the optical axis of the WCR) over the objective lens as well as over time.

The Gershun tube does almost the same, except now the vector product \(\mathbf{E} \times \mathbf{H}\) is integrated over the small exit aperture.

The reader should find it quite instructive to recognize that despite seemingly being quite different, the one natural and four manmade devices shown in Fig. 15 perform the same physical operation of filtering out electromagnetic wavefronts rather than electromagnetic energy currents. Some of these devices can be equipped with two-dimensional pixelated detectors such as a CCD or retina, in which case each pixel has the same functionality as the diaphragm in Fig. 12e besides being an individual photodetector. In the final analysis, all these devices are WCRs, perhaps with the added panoramic capability.

It is remarkable that despite the massive practical use of optical instruments such as those shown in Figs. 12e and 13 for many decades, the fundamental physical principle of the corresponding measurements had not been recognized until quite recently [117]. We will see in the following section that once this principle had been clearly stated, it became possible to formulate the discipline of directional radiometry as a branch of physical optics.

4. Path to physical optics

The tenuous standing of the centuries-old phenomenological disciplines of directional radiometry and radiative transfer with respect to fundamental physics has been recognized for decades [12,21,28,64,109,111,121]. Over the past 50 years many important studies have been published with the goal of developing the requisite microphysical (i.e., back-traceable to the MMEs) foundation for both disciplines (e.g., Refs. [121–139] and references therein). In most cases the heuristic concept of the local polydirectional radiance has been viewed as worth being preserved despite being poorly defined. This has led to attempts to identify a quadratic form in the electric (or electric and magnetic) field vectors that would satisfy the RTE and thereby might be considered a legitimate microphysical proxy for the phenomenological specific intensity. However, those studies have not clarified the issue of the requisite non-negativity and physical measurability of such proxies as well as have not established their relevance to the actual electromagnetic energy transport in turbid media. Furthermore, in many cases the scalar wave equation has been used in lieu of the Maxwell equations. Some studies have been based on artificial mathematical models of a random electromagnetic field wherein certain statistical characteristics of the randomness were prescribed a priori rather than derived from direct solutions of the Maxwell equations for statistically random scattering media. It was thus assumed that the regime of radiative transfer must be attributed to a certain random behavior of the sources of the electromagnetic field rather than to the random behavior of the scattering medium. However, this assumption rules out the case of illumination of a turbid medium by a monochromatic or quasi-monochromatic parallel beam and as such has been shown to be generally incorrect.

As a consequence of various shortcuts, the majority of these publications have achieved only partial success in establishing the microphysical foundation of the KIT and directional radiometry. As recently as in 1995, Leonard Mandel and Emil Wolf remarked (see page 287 of Ref. [109]) that in spite of the long history of radiometry and the theory of radiative energy transfer, their foundations had not been fully clarified.

The final solution of the problem in the case of electromagnetic scattering by a turbid medium has become possible only with the recent realization that one does not need to use the specific intensity as the fundamental point of departure and contemplate it as an actual physical quantity required to possess certain desirable properties as well as satisfy the RTE. Instead of declaring the overarching objective of identifying a microphysical quantity that would allow one to derive the RTE for its own sake, one needs to focus on addressing the following problems that are consistent with the very structure and range of applicability of Maxwell’s electromagnetics as well as have actual and straightforward practical importance:

1. How to evaluate theoretically the time-averaged radiation-energy budget of a macroscopic volume of random particulate medium?
2. Given the widespread practical use of WCRs, how to model theoretically the particular measurement afforded by a WCR and thereby clarify its ability to serve as (i) an energy-budget instrument and/or (ii) an integral part of a
diagnostic technique intended for optical characterization of a particulate medium in a laboratory, in situ, or remote-sensing arrangement?

Indeed, it is the solution of these two specific and well-defined problems that one needs in the final analysis, the hypothetical “angular distribution of the local electromagnetic energy flow” being irrelevant and unnecessary in addition to being unphysical. Both problems have been addressed by directly and self-consistently solving the Maxwell equations, as applied to a turbid medium, in Refs. \[116,140\] (see also Refs. \[117,141\]).

The process of establishing the microphysical foundation of the disciplines of directional radiometry and radiative transfer has been rather long and convoluted. In what follows, I will try to highlight the major milestones of this endeavor. Certain key developments, such as the discovery of the electromagnetic nature of light, have not been rooted specifically in either discipline and have a much broader significance. Yet they have provided the requisite conceptual framework and as such will also be mentioned.

4.1. Maxwell’s electromagnetics

Although the pioneering research by Thomas Young (1773–1829), Augustin Jean Fresnel (1788–1827), and Sir George Gabriel Stokes (1819–1903; see Fig. 16) had established the wave nature of light, it was the existence of a specific plane-wave solution of the MMEs that had led James Clerk Maxwell (1831–1879) to conclude that light consists of electromagnetic waves. Although visible light was the only part of the electromagnetic spectrum known to Maxwell, subsequent research led to the discovery of other parts of the spectrum which now extends from gamma rays to radio waves.

Maxwell’s electromagnetics was the first relativistic field theory and constitutes one of the supreme intellectual achievements in the history of humankind. Ludwig Boltzmann was so enchanted by the beauty and might of the Maxwell equations that he quoted from Johann Goethe’s Faust: “Was it a God who wrote these signs” (see page iii of Ref. [142]). In his magnificent history of mathematical thought from ancient to modern times, Morris Kline wrote: “The most spectacular triumph of the nineteenth century, with an enormous impact on science and technology, was Maxwell’s derivation in 1864 of the laws of electromagnetism” (see page 698 of Ref. [143]).

A meticulous account of the history of classical electromagnetics from the time of Gilbert and Descartes to the relativity theory of Poincaré and Lorentz was given by Sir Edmund Taylor Whittaker [144]. Ref. [145] describes how Maxwell’s ideas, summarized in his famous Treatise [146], were picked up, organized, and reworked mathematically by his immediate followers, most notably by Oliver Heaviside (1850–1925) [147]. An important “byproduct” of Heaviside’s work on electromagnetics was the creation (independently of J. Willard Gibbs) of vector algebra and vector analysis. Jules Henri Poincaré (1854–1912)
introduced the overarching principle of relativity [148] and derived the relativity theory as a direct corollary of the Maxwell equations [149,150].

The famous Poynting theorem quantifying the energy budget of a finite volume element was derived by John Henry Poynting (1852–1914) in 1884 [151] and independently by Heaviside in 1885 [152]. Lorentz [90] gave the first derivation of the MMEs from classical electromagnetics. His work was refined and generalized by de Groot and Suttorp [153] and Robinson [154]. The direct derivation of the MMEs from QED has turned out to be a much more complex problem. It is especially involved if a dielectric medium is lossy, in which case the dielectric needs to be linked to a thermal field reservoir. Recent progress in this direction has been substantial [155–157], but further research is still needed [158].

Classical macroscopic electromagnetics is not the most fundamental physical theory of light–matter interactions. In particular, it ignores the discreteness of matter and operates with continuous sources of fields, which implies that its predictions can fall short in cases where quantum effects are essential. In spite of that, the quantum theory can often be used to determine the “bulk” electromagnetic properties of bodies consisting of very large numbers of atoms [158]. It has been demonstrated that this approach works well when the external electromagnetic field is sufficiently weak and the size of individual particles forming a turbid medium exceeds ~50 Å [159]. Obviously, this result implies a rather wide range of practical applicability of the MMEs.

A clear delineation of the type of field–matter interactions captured by classical macroscopic electromagnetics helps identify specific problems that can be addressed by solving the MMEs. For example, let us consider a cloud of liquid-water droplets illuminated by a parallel quasi-monochromatic beam of light (Fig. 17) and suppose that we need to evaluate the energy budget of a macroscopic volume element ΔV bounded by the closed surface ΔS. According to the Poynting theorem, the net average rate at which electromagnetic energy enters this volume element is given by the surface integral

\[
\langle W_{\Delta S} \rangle = - \int_{\Delta S} d^2r \langle \mathbf{S}(\mathbf{r}, t) \rangle \cdot \hat{n}(\mathbf{r}) \geq 0.
\]  

where \( \langle \cdots \rangle \) denotes averaging over a sufficiently long period of time and the unit vector \( \hat{n}(\mathbf{r}) \) is directed along the local outward normal to the boundary. The meaning of the Poynting theorem is straightforward if \( \langle W_{\Delta S} \rangle = 0 \), in which case the incoming energy is balanced by the outgoing energy and the particulate matter inside ΔV is not affected by the electromagnetic radiation. However, if \( \langle W_{\Delta S} \rangle > 0 \) and ΔV contains no free charges then the Poynting theorem implies that there is a continuous accumulation of electromagnetic energy inside ΔV, which is physically unrealistic. This result exposes an inherent weakness of the MMEs rooted in their inability to describe nonlinear field–matter interactions. A conventional “patch” used to circumvent this issue is the postulate that if \( \langle W_{\Delta S} \rangle > 0 \) then the excess electromagnetic energy is transformed into other forms of energy (e.g., heat) via physical mechanisms not specifically described by classical macroscopic electromagnetics.

Another type of practical problems that can be addressed in the framework of classical macroscopic electromagnetics has to do with the fact that instruments such as WCRs are capable of measuring various manifestations of electromagnetic energy flow. The signal measured by these instruments can carry imbedded information on the physical properties of the scattering medium. To extract this information from the signal, one must (i) have a clear understanding of the physical nature of the measurements and (ii) be able to solve the inverse problem of identifying the physical model of a particulate medium that provides the best fit of theoretical computations of electromagnetic scattering to the measurement data. Of course, an integral part of solving the inverse problem is solving the direct problem, i.e., finding an accurate solution of the MMEs for a given model of particulate medium and a specific type of illumination.

4.2. Electromagnetic scattering

To evaluate theoretically the time-averaged radiation energy budget of a macroscopic volume of particulate medium or to model theoretically the time-averaged reading of a WCR, one must be able to address the following two problems:

- determine the instantaneous electric and magnetic fields by solving the MMEs for a fixed multi-particle configuration and
- average the Poynting vector and relevant optical observables over a sufficiently long period of time during which particle positions and microphysical states are allowed to randomly change.

The microphysical state of a particle is characterized by its size, morphology (including the potentially inhomogeneous distribution of the relative refractive index), and orientation.

To solve the first problem, it is usually assumed that the fixed N-particle group is imbedded in a homogeneous and nonabsorbing infinite host medium. Furthermore, it is assumed that over time intervals much longer than \( 2\pi/\omega \), the time dependence of the electric and magnetic fields everywhere in space is harmonic and described, in the complex-field representation, by the simple complex exponential \( \exp(-i\omega t) \), where \( \omega \) is the angular frequency and \( i = (-1)^{1/2} \). In other words, it is assumed that the complex electric and magnetic fields can be factorized as

\[
\mathbf{E}(\mathbf{r}, t) = \exp(-i\omega t) \mathbf{E}(\mathbf{r}) \quad \text{and} \quad \mathbf{H}(\mathbf{r}, t) = \exp(-i\omega t) \mathbf{H}(\mathbf{r}),
\]

respectively, while the actual real-valued fields are obtained by taking the real part of the respective complex fields:

\[
\mathbf{E}(\mathbf{r}, t) = \text{Re} \mathbf{E}(\mathbf{r}, t) \quad \text{and} \quad \mathbf{H}(\mathbf{r}, t) = \text{Re} \mathbf{H}(\mathbf{r}, t).
\]

The frequency-domain monochromatic Maxwell curl equations for the time-independent electric and magnetic field amplitudes \( \mathbf{E}(\mathbf{r}) \) and \( \mathbf{H}(\mathbf{r}) \) are as follows:

\[
\begin{align*}
\nabla \times \mathbf{E}(\mathbf{r}) &= -i\omega \mu_0 \mathbf{H}(\mathbf{r}) \quad \forall \mathbf{r} \in \Omega_{\text{EXH}}, \\
\nabla \times \mathbf{H}(\mathbf{r}) &= -\frac{i}{\omega \varepsilon_0} \mathbf{E}(\mathbf{r})
\end{align*}
\]  

\[ \mathbf{r} \in \Omega_{\text{EXT}}. \]  

(13)
\[ \nabla \times \mathbf{E}(\mathbf{r}) = \frac{i \omega \mu_0}{\varepsilon_1} \mathbf{H}(\mathbf{r}) \]
\[ \nabla \times \mathbf{H}(\mathbf{r}) = -i \omega \varepsilon_2(\mathbf{r}, \omega) \mathbf{E}(\mathbf{r}) \quad \mathbf{r} \in V_{\text{INT}}. \]  

In these equations, \( V_{\text{INT}} \) is the cumulative “interior” volume occupied by the \( N \) particles (Fig. 18):

\[ V_{\text{INT}} = \bigcup_{i=1}^{N} V_i. \]

where \( V_i \) is the volume occupied by the \( i \)th particle; \( V_{\text{EXT}} \) is the infinite exterior region such that \( V_{\text{INT}} \cup V_{\text{EXT}} = \mathbb{R}^3 \), where \( \mathbb{R}^3 \) denotes the entire three-dimensional space; the host medium and the scattering object are assumed to be nonmagnetic; \( \mu_0 \) is the permeability of a vacuum; \( \varepsilon_1 \) is the real-valued electric permittivity of the host medium; and \( \varepsilon_2(\mathbf{r}, \omega) \) is the (potentially coordinate-dependent) complex permittivity of the \( N \)-particle object.
The corresponding boundary conditions read

\[ \mathbf{n} \times (\mathbf{E}_1(r) - \mathbf{E}_2(r)) = \mathbf{0}, \quad \mathbf{n} \times (\mathbf{H}_1(r) - \mathbf{H}_2(r)) = \mathbf{0}, \quad r \in S_{\text{INT}}, \]

where \( \mathbf{0} \) is a zero vector, the subscripts 1 and 2 correspond to the exterior and interior sides of the composite boundary \( S_{\text{INT}} \) of the \( N \)-particle object, respectively, and \( \mathbf{n} \) is the local outward normal to \( S_{\text{INT}} \). In agreement with Eq. (15), \( S_{\text{INT}} \) is the union of the closed surfaces of the \( N \) particles

\[ S_{\text{INT}} = \bigcup_{i=1}^{N} S_i. \]

Let us assume that the total field \( (\mathbf{E}(r), \mathbf{H}(r)) \) everywhere in space can be represented by a vector superposition of a plane-wave component (traditionally called the “incident field”); superscript “inc”) propagating in the direction of the unit vector \( \mathbf{n}^{\text{inc}} \) and a “scattered field” (superscript “sca”):

\[ \mathbf{E}(r) = \mathbf{E}_0^{\text{inc}} \exp \left( \frac{\mu_0}{i} \mathbf{n}^{\text{inc}} \cdot \mathbf{r} \right) + \mathbf{E}^{\text{sca}}, \]
\[ \mathbf{H}(r) = \frac{\mathbf{E}_0^{\text{inc}}}{\mathbf{\mu}_0} \times \mathbf{n}^{\text{inc}} \exp \left( \frac{\mu_0}{i} \mathbf{n}^{\text{inc}} \cdot \mathbf{r} \right) + \mathbf{H}^{\text{sca}}, \]

where \( k_1 = \omega (\epsilon_1 \mu_0)^{1/2} \) is the wave number of the exterior region. In addition, we postulate that the scattered field satisfies the following condition at infinity:

\[ \lim_{r \to \infty} \left\{ \sqrt{\mu_0} \mathbf{r} \times \mathbf{H}^{\text{sca}}(r) + r \sqrt{\epsilon_1} \mathbf{E}^{\text{sca}}(r) \right\} = \mathbf{0}, \]

where \( r = \mathbf{r} \) is the distance from the observation point to the fixed origin \( O \) located near the “geometrical center” of the \( N \)-particle group (Fig. 18).

The curl Eqs. (13) and (14) supplemented by the boundary conditions (16), the decomposition (18), and the asymptotic condition (19) constitute the standard electromagnetic scattering problem for plane-wave illumination.

The above formulation of the standard electromagnetic scattering problem is hardly useful unless this problem has a solution and unless this solution is unique. To the best of my knowledge, the existence and the uniqueness of the solution of the standard scattering problem have been proven only in some particular cases. However, the thorough analysis by Claus Müller [160] demonstrates the fundamental importance of imposing both the boundary conditions and the asymptotic condition at infinity. It is, therefore, reasonable to assume that the standard scattering problem does have a solution, this solution being unique.

The pioneers of the theory of electromagnetic scattering by particles John William Strutt, Lord Rayleigh (1842–1919), Ludvig Valentin Lorenz (1829–1891), and Gustav Mie (1868–1957) considered it to be intuitively obvious that the “physically relevant” solution of the MMEs must involve the scattered field in the form of an outgoing spherical wave. Arnold Sommerfeld (1868–1951) was the first to realize in 1912 that the very uniqueness of the solution of an open-space scattering problem depends on the explicit postulation of an asymptotic condition at infinity [161]. The history of this so-called Sommerfeld radiation condition as well as its extensions and modifications are described in Ref. [162]. In the specific case of three-dimensional electromagnetic scattering by finite objects, the Sommerfeld radiation condition takes the form of Eq. (19) and is traditionally called the Silver–Müller radiation condition [160,163].

While the standard scattering problem is intentionally formulated for the incident field in the form of a plane wave.
electromagnetic wave, its actual range of relevance is much wider since the MMEs, the boundary conditions, and the radiation condition are all linear in the electric and magnetic fields. This implies that if \{\mathbf{E}, \mathbf{H}\} and \{\mathbf{E}'', \mathbf{H}''\} are the solutions of the standard problem corresponding to the incident plane waves \{\mathbf{E}_i, \mathbf{H}_i\} and \{\mathbf{E}_i'', \mathbf{H}_i''\}, respectively, then \{\mathbf{E} + \mathbf{E}'', \mathbf{H} + \mathbf{H}''\} is the unique solution of the boundary-value scattering problem for the incident field given by \{\mathbf{E}_i, \mathbf{H}_i\}. Hence solutions of the standard scattering problem can be used to obtain the solution of a more general scattering problem provided that the corresponding incident field can be expanded in plane electromagnetic waves.

4.3. Volume integral equation and Lippmann–Schwinger equation

Although the standard scattering problem is stated as a boundary-value problem for the differential frequency-domain MMEs, it is often convenient to invoke an equivalent integral-equation formulation. Specifically, the so-called volume integral equation (VIE) expresses the total electric field everywhere in space in terms of the incident electric field and the total electric field in the interior volume of the N-particle group. The VIE is a straightforward consequence of Eqs. (13)–(19) and reads

$$\mathbf{E}(\mathbf{r}) = \mathbf{E}_i(\mathbf{r}) + k_i^2 \int_{\text{INT}} d^3r' \left\{ m^2(\mathbf{r}' - 1) \mathbf{G}(\mathbf{r}, \mathbf{r}') \cdot \mathbf{E}(\mathbf{r}') \right\}, \quad \mathbf{r} \in \mathbb{R}^3,$$

(20)

where \(m(\mathbf{r}) = \kappa_0(\mathbf{r})/k_i\) is the refractive index of the interior relative to that of the exterior, \(k_0(\mathbf{r}) = \omega \mu_0(\mathbf{r}) \cdot \omega \mu_0(\mathbf{r}) \cdot 1/2\) is the wave number of the interior region,

$$\mathbf{G}(\mathbf{r}, \mathbf{r}') = \left( \mathbf{I} + \frac{1}{k_i^2} \mathbf{V} \otimes \mathbf{V} \right) \exp(i \mathbf{k}_i(\mathbf{r} - \mathbf{r}')), \quad \mathbf{r}, \mathbf{r}' \in \mathbb{R}^3,$$

(21)

is the free-space dyadic Green's function, \(\mathbf{I}\) is the identity dyadic, and \(\otimes\) is the dyadic product sign. A major advantage of the VIE is that the boundary conditions (16) and the radiation condition (19) are built into it rather than serve as additional constraints imposed on the physically relevant solution of the differential frequency-domain MMEs. The electric field found by solving the VIE can be substituted in the second formulas of Eqs. (13) and (14) in order to find the magnetic field everywhere in space.

The dyadic Green's function was introduced in 1950 by Harold Levine and Julian Schwinger [164]. It belongs to the general class of functions first studied by the self-taught English mathematician (and professional miller) George Green (1793–1841) in his initially obscure, but now famous Essay of 1828 privately printed at Green’s own expense in the form of a booklet [165]. Typically, a Green’s function is an integral kernel that can be used to solve an inhomogeneous differential equation defined on a domain, with specified boundary conditions. The Essay was enthusiastically popularized by William Thomson, Lord Kelvin who arranged its re-publication in *Jahrb. für die reine und angewandte Mathematik* (also known as *Crelle’s Journal*) in 1850–1854. Green’s approach was further developed into an efficient mathematical tool by Oliver Heaviside.

Note that the majority of derivations of the VIE published in monographs and journal papers, including the original derivation by David Saxon (1920–2005) in 1955 [166], do not invoke explicitly the boundary conditions (16) as the prerequisite of the uniqueness of solution of the standard scattering problem. As such, they are applicable only to “fuzzy” scattering objects without distinct boundaries. This means that the refractive index is implicitly assumed to depend on spatial coordinates continuously throughout the entire space rather than being discontinuous across particle surfaces. A complete derivation explicitly relying on Eq. (16) and thereby applicable to scattering objects with distinct boundaries is given in Refs. [140,167].

The VIE can be perceived as the very embodiment of the concept of electromagnetic scattering. Indeed, it shows that in the absence of the scattering object, i.e., when \(m(\mathbf{r}) = 1\), the total field is identically equal to the incident field. The presence of the object changes the total field, which means that the scattered field can be defined as the difference between the total fields in the presence and in the absence of the object. Furthermore, the VIE makes it obvious that the incident field is not modified by the presence of the object and, contrary to the widespread misconception, is not “transformed into the scattered field” [168].

It is convenient to express the scattered field mathematically in terms of the incident field as follows:

$$\mathbf{E}^{\text{sc}}(\mathbf{r}) = \int_{V_{\text{INT}}} d^3r' \, \mathbf{G}(\mathbf{r}, \mathbf{r}') \cdot \mathbf{E}(\mathbf{r}), \quad \mathbf{r} \in \mathbb{R}^3,$$

(22)

where \(\mathbf{G}\) is the so-called dyadic transition operator (DTO). Substituting Eq. (22) in Eq. (20) yields the following integral equation for \(\mathbf{G}\):

$$\mathbf{G}(\mathbf{r}, \mathbf{r}') = k_i^2[1/m(\mathbf{r}') - 1] \mathbf{E}(\mathbf{r}') \cdot \mathbf{I} + k_i^2[1/m(\mathbf{r}') - 1] \int_{V_{\text{INT}}} d^3r'' \, \mathbf{G}(\mathbf{r}, \mathbf{r}'') \cdot \mathbf{E}(\mathbf{r}'').$$

(23)

Equations of this type were introduced in 1950 by Bernhard A. Lippmann and Julian Schwinger in the quantum theory of scattering [169] and are traditionally called Lippmann–Schwinger equations [170,171]. The first derivation of Eq. (23) in the electromagnetic-scattering case was given by Leung Tsang and Jin Au Kong in 1980 [172] (see also Refs. [173,174]).

A fundamental property of the DTO is that it is independent of the incident field and is a function of the scattering object only: it is fully defined by the distribution of the relative refractive index throughout \(V_{\text{INT}}\). It is this property that has made the concept of the DTO central to the theory of electromagnetic scattering by multi-particle groups. In particular, as discussed in the following subsection, it was used to derive the so-called Foldy equations (FEs) that eventually served as a natural precursor to the microphysical theory of radiative transfer in sparse particulate media.
4.4. Foldy equations

If the scattering object is a multi-particle group, then it is very convenient to formally represent the total scattered field as a vector superposition of the “partial” fields contributed by the individual particles. In the case of an arbitrary N-particle group, this cannot be done easily using the differential frequency-domain Maxwell equations, but is quite straightforward in the framework of the VIE formalism. Indeed, let us first re-write the VIE as follows:

\[ \tilde{E}(r) = E^{\text{inc}}(r) + \sum_{i=1}^{N} U_i(r), \quad r \in \mathbb{R}^3, \]  

(24)

where the integration is performed over the entire space, the potential function \( U(r) \) is given by

\[ U(r) = \sum_{i=1}^{N} U_i(r), \quad r \in \mathbb{R}^3, \]

(25)

and \( U_i(r) \) is the potential function of the \( i \)-th particle. The latter is given by

\[ U_i(r) = \begin{cases} 0, & r \notin V_i, \\ k_i^2[m_i^2(r) - 1], & r \in V_i, \end{cases} \]

(26)

where

\[ m_i(r) = k_0(r)/k_1 \]

is the refractive index of particle \( i \) relative to that of the host medium. All position vectors originate at the common origin \( O \) of the laboratory coordinate system (Fig. 18). It can then be shown that the total electric field everywhere in space is given by the following expression:

\[ \tilde{E}(r) = E^{\text{inc}}(r) + \sum_{i=1}^{N} \int_{V_i} d^3 \tilde{r} \tilde{G}(r, \tilde{r}) \sum_{j \neq i} \tilde{T}_j(r, \tilde{r}) \cdot \tilde{E}_j(r), \quad r \in \mathbb{R}^3, \]

(28)

where the total electric field \( \tilde{E}(r) \) “exciting” particle \( i \) is given by

\[ \tilde{E}_i(r) = E^{\text{inc}}(r) + \sum_{j \neq i} \tilde{E}_j^{\text{exc}}(r), \]

(29)

the \( \tilde{E}_j^{\text{exc}}(r) \) are partial exciting fields given by

\[ \tilde{E}_j^{\text{exc}}(r) = \int_{V_j} d^3 \tilde{r} \tilde{G}(r, \tilde{r}) \cdot \tilde{T}_j(r, \tilde{r}) \cdot \tilde{E}_j(r), \quad r \in V_j, \]

(30)

and \( \tilde{T}_i \) is the \( i \)-th-particle DTO defined with respect to the laboratory coordinate system; it satisfies the following Lippmann–Schwinger equation:

\[ \tilde{T}_i(r, \tilde{r}) = U_i(r) \delta(r - \tilde{r}) + \int_{V_i} d^3 \tilde{r} \tilde{G}(r, \tilde{r}) \cdot \tilde{T}_i(r, \tilde{r}) \cdot \tilde{E}_j(r), \quad \tilde{r} \in V_i. \]

(31)

Importantly, each \( \tilde{T}_i \) is computed individually, as if all the other particles did not exist.

A simplified version of Eqs. (28)–(31) was proposed heuristically in 1945 by Leslie L. Foldy (1919–2001) [175] to describe scattering of scalar waves by a system of widely separated isotropic point centers. Various applications of Foldy’s approach were discussed in 1951 by Melvin Lax [176]. A rigorous derivation of the quantum-mechanical version of the FE from the Schrödinger equation was given in 1953 by Kenneth M. Watson [177] (see also Section 11.3 of Ref. [170]). The case of electromagnetic scattering by a group of widely separated electrons was considered by Watson in 1969 [178]. The general electromagnetic FE of the form of Eqs. (28)–(31) were derived by Tsang and Kong in 1980 [172] (see also Ref. [173] in which the electromagnetic FE were derived using Watson’s approach).

The FE of Eq. (28)–(31) (also known as Foldy–Lax equations [136,179]) are mathematically equivalent to the VIE (20) and hardly offer computational advantages. However, there are two factors which make them important. First, they allow one to introduce the Neumann expansion of the total field playing a key role in the microphysical theory of radiative transfer. Second, as discussed in the following subsection, they can be used to incorporate the notion of the single-particle far field in the computation of the near field of a multi-particle group.

Let us first define the \( i \)-th potential dyadic centered at the origin of the laboratory reference frame according to

\[ \tilde{U}_i(r, \tilde{r}) = U_i(r) \delta(r - \tilde{r}) \tilde{T}_i \]

(32)

and introduce the following operator notation:

\[ \tilde{U} = \sum_{i=1}^{N} \tilde{U}_i, \]

(33)

\[ B_E = \int d^3 \tilde{r} \tilde{B}(r, \tilde{r}) \cdot \tilde{E}(r). \]

(34)

Iterating Eqs. (29) and (30) yields

\[ E = E^{\text{inc}} + \sum_{j \neq i} \tilde{C}_j^{\text{inc}} E^{\text{exc}} + \sum_{j \neq i} \sum_{k \neq j} \tilde{C}_j^{\text{inc}} \tilde{C}_k^{\text{inc}} E^{\text{exc}} + \ldots, \]

(35)

whereas the substitution of Eq. (35) in Eq. (28) results in the following expansion, traditionally called in mathematics the Neumann series:

\[ E = E^{\text{inc}} + \sum_{i=1}^{N} \tilde{C}_i^{\text{inc}} E^{\text{exc}} + \sum_{i=1}^{N} \sum_{j \neq i} \tilde{C}_j^{\text{inc}} \tilde{C}_k^{\text{inc}} E^{\text{exc}} + \ldots \]

(36)

It is easily seen that the Neumann series is fundamentally based on the fact that \( \tilde{T}_i \) for each \( i \) is an individual property of the \( i \)-th particle computed as if this particle were alone rather than a member of the group. As a consequence, there has been a tendency to characterize the Neumann expansion (36) as describing “multiple scattering” by the \( N \)-particle group. It is important to recognize, however, that although the notion of multiple scattering, as embodied by Eq. (36), can be a useful mathematical abstraction, it is not a real physical phenomenon wherein the incident light is scattered sequentially by one, two, three or more particles before reaching the observation point [168,180]. In other words, one can speak of multi-particle sequences contributing the various terms on the right-hand side of Eq. (36), but not of actual “multiple-scattering paths” or “multiple-scattering trajectories".
The concept of the Neumann expansion (also known as the Liouville–Neumann or Liouville–Neumann–Volterra series) was introduced by the French mathematician Joseph Liouville (1809–1882) in 1837 and by the German mathematician Carl Gottfried Neumann (1832–1925) thirty years later. The Italian mathematician Vito Volterra (1860–1940) used this expansion in his general theory of integral equations as a technique for solving Fredholm and Volterra equations of the second kind, wherein the unknown function is expanded in a power series in terms of so-called iterated kernels. The method is applicable whenever the series converges [181,182]. In quantum theory of particle scattering, an expansion of the type (36) is often called the Born series [170,171].

4.5. Far-field Foldy equations

The reader should recognize that the formulation of electromagnetic scattering in Sections 4.1–4.4 has been quite general. Although we chose to speak of the scattering object in the form of a fixed group of $N$ “distinct” non-overlapping particles, Eqs. (13)–(36) apply to any morphology of the scattering object as long as the entire object is finite. For example, we could consider a single ellipsoid and subdivide it into an arbitrary set of $N$ non-overlapping volume elements, in which case the FEs would still hold. This demonstrates again that the concept of multiple frequency-domain scattering, as represented by the Neumann expansion (36), is nothing more than a mathematical abstraction [180].

As already mentioned, the FEs offer no computational advantages over the VIE in the case of an arbitrary fixed scattering object. However, they become indispensable if the scattering object is a large, sparse, random cloud of particles, in which case the Neumann expansion (36) helps accommodate a very straightforward analytical fashion two main morphological traits of the object: large particle-to-particle distances and statistically uncorrelated particle positions.

Specifically, let us first assume that:

- the particles forming the group are separated widely enough that each of them is located in the far zones of all the other particles, and
- the observation point is located in the far zone of any particle in the group (but not necessarily in the far zone of the entire group).

As a consequence, each partial exciting field $E_{exc}^{inc}(r)$ in Eq. (29) becomes an outgoing spherical wavelet centered at particle $j$, while the integral FEs are converted into algebraic far-field FEs. The latter imply that the total field at any observation point located sufficiently far from any particle in the sparse multi-particle group is the superposition of the incident plane wave and $N$ partial spherical wavelets contributed by the $N$ particles. The observation point is not required to be in the far zone of the entire group. It can be anywhere in space as long as it resides in the far zones of all the particles forming the group.

It is important to recognize that even though each component of the cumulative scattered field given by the far-field FEs is a transverse electromagnetic wave, the scattered field itself is not, in general, a transverse electromagnetic wave. In other words, the far-field FEs describe the far field of the multi-particle group despite the underlying assumption that the observation point is located in the far zone of any constituent particle. The cumulative scattered field becomes a transverse electromagnetic wave only if the observation point is located in the far zone of the entire group.

The corresponding far-field Neumann expansion of the total electric field also becomes purely algebraic and takes the form

$$E = E_{inc} + \sum_{i=1}^{N} B_{rij} \cdot E_{ij}^{inc} + \sum_{i=1}^{N} \sum_{j=1}^{N} B_{rij} \cdot B_{gij} \cdot E_{ij}^{inc} + \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{m=1}^{N} B_{rij} \cdot B_{gij} \cdot B_{gjm} \cdot E_{ij}^{inc} + \cdots,$$

(37)

where each dyadic $B_{gij}$ describes far-field scattering by the $j$th particle. It is Eq. (37) that represents the starting point in the development of the microphysical theory of radiative transfer.

The far-field electromagnetic FEs were introduced heuristically by Anatoly Borovoi for a group of identical spherical particles in 1966 [122]. The first rigorous derivation from the general electromagnetic FEs (28)–(31) appeared in 1984 [173] (see also Refs. [137,183]).

4.6. The Twersky approximation

The terms with $j=i$ and $l=j$ in the triple summation on the right-hand side of the far-field Neumann expansion (37) are excluded, but the terms with $i=l$ are not. Therefore, we can decompose this summation as follows:

$$\sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{l=1}^{N} B_{rij} \cdot B_{gij} \cdot B_{gjm} \cdot E_{ij}^{inc}$$

$$= \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{l=1}^{N} B_{rij} \cdot B_{gij} \cdot B_{gjm} \cdot E_{ij}^{inc}$$

$$+ \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{l=1}^{N} B_{rij} \cdot B_{gij} \cdot B_{gjm} \cdot E_{ij}^{inc}.$$

(38)

Higher-order summations in Eq. (37) can be decomposed similarly. Thus, the total field at an observation point $r$ is composed of the incident field, one-particle contributions, and contributions from multi-particle sequences that can be divided into two groups. The first one includes all the terms contributed by self-avoiding multi-particle sequences, whereas the second group includes all the terms corresponding to multi-particle sequences that involve a particle more than once. The approximation introduced in 1964 by Victor Twersky [184] (for the case
of scattering of scalar waves) results in a drastic simplification by neglecting the terms belonging to the second group and retaining only the terms from the first group:

$$
\mathbf{E} \approx \mathbf{E}_{\text{inc}} + \sum_{i=1}^{N} \sum_{j=1}^{N} B_{ij} \cdot \mathbf{E}_{i}^{\text{inc}} + \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{l=1}^{N} B_{ijl} \cdot B_{jl0} \cdot \mathbf{E}_{i}^{\text{inc}} + \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{l=1}^{N} \sum_{m=1}^{N} B_{ijlm} \cdot B_{l0m} \cdot B_{m00} \cdot \mathbf{E}_{m}^{\text{inc}} + \ldots. \quad (39)
$$

It can be shown indeed that in the limit $N \to \infty$ the Twersky approximation accounts for the overwhelming majority of multi-particle sequences, which implies that it can be expected to yield rather accurate results provided that the number of particles in the group is sufficiently large.

4.7. The Poynting–Stokes tensor and dyadic correlation function

The Twersky expansion (39) coupled with the first equation of Eq. (13) can be used to write an extended expression for the time average of the complex Poynting vector $\mathbf{S}(\mathbf{r}) = \frac{1}{2} \mathbf{E}(\mathbf{r}) \times \mathbf{H}^*(\mathbf{r})$, where the asterisk denotes a complex-conjugate value. However, doing that does not enable one to contract the resulting infinite set of nested summations and obtain a closed-form equation amenable to efficient analytical or numerical solution. The ultimate reason for that is the following inescapable fact: different combinations of electric and magnetic field vectors can yield the same Poynting vector. This means that forming the vector product of the electric and magnetic field vectors results in a quantity that does not carry unique information about the participating fields. As a consequence, the Poynting vector cannot be used to describe the phenomenon of electromagnetic scattering by, for example, expressing the Poynting vector of the total field in that of the incident field.

It is also customary to describe electromagnetic scattering in the far zone of a finite object in terms of the $4 \times 4$ Stokes phase matrix using the 4-element Stokes column vector as the primary descriptor of polarization. However, this formalism can be applied only to transverse (e.g., plane or spherical) electromagnetic waves, whereas the total electromagnetic field in the near zone of any object (e.g., at any observation point inside a multi-particle group) is never a transverse wave.

It was shown by the author in 2010 [116] (see also Ref. [140]) that instead of attempting to calculate directly the time-averaged Poynting vector or Stokes column vector, one should first calculate the time average of the Poynting–Stokes tensor (PST) defined as the dyadic product of the electric and magnetic field vectors:

$$
\mathbf{P}(\mathbf{r}) = \frac{1}{2} \mathbf{H}(\mathbf{r}) \otimes \mathbf{E}^*(\mathbf{r}). \quad (40)
$$

It is fundamentally important that by its very construct, the PST is applicable to an arbitrary electromagnetic field (including the near field) and thus can be used to find both the Poynting vector and, whenever applicable, relevant optical observables such as the Stokes parameters.

The PST involves both the electric and the magnetic field at the observation point $\mathbf{r}$. It is sometimes convenient to have an alternative representation involving only the electric field. It can be verified easily that in the case of nonmagnetic materials, Eq. (40) everywhere in space can be written in the form

$$
\mathbf{P}(\mathbf{r}) = \frac{1}{2 \mu_0 \varepsilon_0} \left[ \nabla \times \mathbf{C}(\mathbf{r}, \mathbf{r}') \right]_{\mathbf{r} = \mathbf{r}'}. \quad (41)
$$

where

$$
\mathbf{C}(\mathbf{r}, \mathbf{r}') = \mathbf{E}(\mathbf{r}) \otimes \mathbf{E}^*(\mathbf{r}). \quad (42)
$$

is the so-called dyadic correlation function involving the electric field at two different points in space. The subscript $\mathbf{r}'$ means that the $\nabla$ operator acts only on $\mathbf{E}(\mathbf{r}')$.

Using the Twersky expansion (39), we can formulate the Twersky approximation for the dyadic correlation function diagrammatically according to Fig. 19. The different terms entering the expanded expression inside the angular brackets on the right-hand side of this equation can be classified using the notation introduced in Fig. 20a. In this particular case, the upper and lower multi-particle sequences involve different particles. However, the two multi-particle sequences can involve one or more common particles, as shown in Fig. 20c–f by using the dashed connectors. Moreover, if the number of common particles in a diagram is two or more, then they can enter the upper and lower sequences in the same order, as in Fig. 20d, or in the reverse order, as in Fig. 20e. Finally, Fig. 20f gives an example of a mixed diagram wherein two common particles appear in the same order while two other common particles appear in the reverse order.

4.8. Ergodicity

So far we have been considering electromagnetic scattering by a fixed configuration of particles. Let us now assume that the sparse N-particle group varies in time, but does it slowly enough that any significant (i.e., modifying the solution of the MMEs) changes of the group occur over time intervals $\tau_0$ much longer than the period of time-harmonic oscillations of the electromagnetic field. Then the time average of the dyadic correlation function is formally defined by

$$
\langle \mathbf{C}(\mathbf{r}', \mathbf{r}, t) \rangle = \frac{1}{T} \int_{t-(T/2)}^{t+(T/2)} dt \mathbf{C}(\mathbf{r}', \mathbf{r}, t'). \quad (43)
$$

where the time interval $T$ is much longer than $\tau_0$. During this time interval the variable N-particle group goes through an infinite sequence of evolving discrete states governed by relevant physical and chemical processes. Therefore, the right-hand side of Eq. (43) must be evaluated by: (i) tracing the temporal evolution of the physical state of the entire
group (i.e., the complete set of particle coordinates and microphysical states), and (ii) computing $\bar{C}(\bf{r}', \bf{r}, t)$ for a sufficiently representative set of moments $t' \in [t-T/2, t+T/2]$.

Although a computer implementation of this averaging procedure is not totally inconceivable, the direct use of the definition (43) in analytical derivations – such as the calculation of the time-averaged dyadic correlation function for a cloud of particles – typically leads to insurmountable technical difficulties. A much more practicable approach is based on the assumption that the $N$-particle group is statistically random and sufficiently variable in time while the time interval $T$ is sufficiently long that averaging $\bar{C}(\bf{r'}, \bf{r}, t)$ over this interval is essentially equivalent to averaging $\bar{C}(\bf{r}, \bf{r}, \bf{r}, \psi)$ over an appropriate analytical probability distribution of the physical state $\psi$ of the group. In other words, it is assumed that averaging over time for one specific realization of the random scattering process is equivalent to ensemble averaging. The equivalence of the time and ensemble averages is called ergodicity.

The ergodic hypothesis was introduced by James Clerk Maxwell and Ludwig Boltzmann (1844–1906) as a basic underlying principle of statistical mechanics and kinetic theory. The mathematical foundation of the ergodic theory, its relation to the famous Poincaré recurrence theorem [185], and applications to statistical physics are discussed in Refs. [186–188].

4.9. Time-averaged Poynting vector

According to the preceding subsection, instead of calculating the average $\langle \langle \bar{C}(\bf{r}, \bf{r}, t) \rangle \rangle$, one can calculate the average $\langle \langle \bar{C}(\bf{r}, \bf{r}, \bf{R}, \xi) \rangle \rangle_{\bf{R}}$, where $\bf{R}$ denotes the complete set of particle coordinates and $\xi$ denotes the complete set of particle microphysical states. Although this problem remains very complex in general, it becomes manageable upon further assuming that:

1. The position and microphysical state of each particle are statistically independent of each other and of those of all the other particles.
2. The microphysical states of all the particles have the same statistical characteristics.
3. The spatial distribution of the particles throughout the medium is completely random and statistically uniform.
4. All diagrams with crossing connectors in the diagrammatic expansion of the dyadic correlation function (cf. Figs. 19 and 20) can be ignored. This is the essence of the so-called ladder approximation [125].

All intermediate steps of the subsequent analytical derivation are detailed in Ref. [140]. Each $\bar{B}$-dyadic in the Twersky expansion (39) includes a complex exponential factor of the type $\exp(ikqR)$, where $R$ can be the distance between two particles or the distance from a particle to the observation point. Analytical averaging of these rapidly oscillating exponentials over uncorrelated and random particle positions leads to dramatic simplifications and as such is at the very heart of the microphysical RTT. The final expression for the time-averaged Poynting vector at an internal observation point $\bf{r}$ is as follows:

$$\langle \langle \bar{S}(\bf{r}, t) \rangle \rangle = \text{Re} \langle \bar{S}(\bf{r}, \bf{R}, \xi) \rangle_{\bf{R}} = \int_{4\pi} d\bf{q} \bar{q} \bar{I}(\bf{r}, \bf{q}).$$

(44)

where the integration is performed over all directions of the unit vector $\bf{q}$ and $\bar{I}(\bf{r}, \bf{q})$ is the first element of the 4-element column

$$\bar{I}(\bf{r}, \bf{q}) = \begin{bmatrix} \bar{I}(\bf{r}, \bf{q}) \\ \bar{Q}(\bf{r}, \bf{q}) \\ \bar{U}(\bf{r}, \bf{q}) \\ \bar{V}(\bf{r}, \bf{q}) \end{bmatrix}. \tag{45}$$

The latter is the solution of the following matrix integro-differential equation:

$$\dot{\bf{q}} \cdot \mathbf{\hat{V}}(\bf{r}, \bf{q}) = -n_0(\bf{K}(\bf{q}, \xi))_0 \bar{I}(\bf{r}, \bf{q}) + n_0 \int_{4\pi} d\bf{q}' \langle \bf{Z}(\bf{q}, \bf{q}', \xi) \rangle \cdot \bar{I}(\bf{r}, \bf{q}). \tag{46}$$

where $n_0$ is the average number of particles per unit volume, $(\bf{K}(\bf{q}, \xi))_0$ is the $4 \times 4$ real-valued single-particle extinction matrix averaged over the microphysical states of all the $N$ particles, and $(\bf{Z}(\bf{q}, \bf{q}', \xi))_0$ is the $4 \times 4$ real-valued single-particle phase matrix, also averaged over the microphysical states of all the $N$ particles. The elements of the matrices $\bf{K}(\bf{q}, \xi)$ and $\bf{Z}(\bf{q}, \bf{q}', \xi)$ are expressed in the elements of the
so-called $2 \times 2$ amplitude scattering matrix describing far-field scattering of a plane electromagnetic wave by an isolated particle with microphysical characteristics described by the state $\xi$ [139,140].

I provide no details of the derivation leading to Eq. (44) because this derivation contains no new concepts and is a straightforward, albeit nontrivial and instructive, mathematical exercise. It is worth noting, however, that the angular argument $\hat{q}$ of $\tilde{I}(\mathbf{r}, \hat{q})$ (as well as of $\tilde{I}(\mathbf{r}, \hat{q})$) at an internal point $\mathbf{r}$ spans the entire range of directions, $\hat{q} \in 4\pi$, since for any $\hat{q}$ there is always a contribution to $\langle \hat{S}(\mathbf{r}; \mathbf{R}, \hat{q}) \rangle_{\mathbf{R}, \xi}$ from pairs of multi-particle sequences such that the left-most common particle of each pair is located in the direction $-\hat{q}$ relative to $\mathbf{r}$.

4.10. Time-averaged reading of a WCR

Consider now a WCR placed inside the random particulate medium, as shown in Fig. 21. The discussion in Section 3 implies that in the framework of the far-field Neumann expansion (39), this instrument reacts only to partial wavelets generated by multi-particle sequences having their end particles located within the conical acceptance volume $\Delta V_q$ defined by the WCR’s small acceptance solid angle $\Omega_0$. Thus, the reading of the instrument is defined not by the full time-averaged PST, but rather by the corresponding partial PST $\langle \hat{P}(\mathbf{r}, \hat{q}; \Delta V_q) \rangle_{\mathbf{R}, \xi}$, where the subscripts $\mathbf{R}$ and $\xi$ denote averaging over coordinates and states of all the $N$ particles constituting the medium and not just those located inside $\Delta V_q$. $\langle \hat{P}(\mathbf{r}, \hat{q}; \Delta V_q) \rangle_{\mathbf{R}, \xi}$ can be computed by making the standard assumptions invoked previously to calculate $\langle \hat{P}(\mathbf{r}) \rangle_{\mathbf{R}, \xi}$, but also requiring that the end particle of any multi-particle sequence be located inside the acceptance volume $\Delta V_q$. This lengthy yet straightforward computation [140] shows that the reading of a WCR shown in Fig. 21 is given by

$$\langle \langle \text{EM power} (\mathbf{r}, \hat{q}) \rangle \rangle \approx S_0 \int d\hat{q} \langle \tilde{I}(\mathbf{r}, \hat{q}) \rangle, \quad (47)$$

where the unit vector $\hat{q}$ specifies the orientation of the optical axis of the WCR and $S_0$ is the surface area of the objective lens. If the WCR can measure all four Stokes parameters, then its polarized reading per unit time is given by

$$\langle \langle \text{Signal} (\mathbf{r}, \hat{q}) \rangle \rangle \approx S_0 \int d\hat{q} \langle \tilde{I}(\mathbf{r}, \hat{q}) \rangle. \quad (48)$$

The fact that the signal recorded by the WCR can be modeled theoretically by solving the RTE often makes the (WCR, RTE) combination a useful optical-characterization tool. Furthermore, comparison of Eqs. (44) and (47) reveals that a WCR can be used to solve the energy-budget problem experimentally by integrating its reading over the entire range of WCR’s orientations $\hat{q} \in 4\pi$. Of course, to make such optical-characterization and energy-budget applications of WCRs possible, the random particulate medium must possess the specific macro- and microphysical properties discussed earlier.

4.11. Corollaries of the microphysical approach to directional radiometry and radiative transfer

Eqs. (44)–(48) can easily be generalized to include the case of an external observation point [140] and finalize the solution of the two key problems formulated at the beginning of Section 4. The implications of the direct derivation of these formulas from the frequency-domain MMEs are quite profound and can be formulated as follows [140].

1. The derivation of Eqs. (44)–(48) does not need fundamental physical laws other than the MMEs. In particular, the ill-defined concepts of independently scattering particles, collective effects, elementary volume elements, incoherent light rays, and photons as localized particles of light have no relevance whatsoever to the transport of electromagnetic radiation in elastically scattering particulate media. Although Eq. (46) (traditionally called the RTE) has the formal mathematical structure of a kinetic equation describing the transport of point-like particles [189], it follows directly from the electromagnetic wave theory.

2. The 4-element column (45) (traditionally called the specific intensity column vector) does not characterize the instantaneous distribution of the radiation field inside the particulate medium. Instead, it emerges as a result of averaging over a sufficiently long period of time. The minimal averaging time necessary to ensure statistical ergodicity may vary depending on the particulate medium in question, but the following is always true: the longer the averaging time the more accurate the theoretical prediction based on Eqs. (44), (47), and (48). The accumulation of a signal over an extended time interval is often used to improve the
measurement accuracy by suppressing the effect of random noise. However, the situation with Eqs. (44)–(48) is fundamentally different since the very applicability of these formulas relies upon averaging over a sufficiently long period of time.

3. To ensure the applicability of Eqs. (44)–(48), the overall size of the particulate medium must be much greater than the wavelength, the average particle size, and the average distance between two neighboring particles.

4. In the context of Eqs. (44)–(48), each particle with its individual extinction and phase matrices is effectively replaced with a virtual random particle characterized by the extinction and phase matrices obtained by averaging over the microphysical states of all the particles.

5. The RTE is an inherently matrix equation involving all four elements of the specific intensity column vector. Its frequently used scalar version is obtained by artificially replacing $\mathbf{i}(\mathbf{r}, \mathbf{q})$ with its first element $i(\mathbf{r}, \mathbf{q})$ (traditionally called the specific intensity or radiance) and the extinction and phase matrices with their respective (1,1) elements. As such, the scalar approximation has no fundamental physical justification.

6. All four elements of the specific intensity column vector are real-valued quantities. Furthermore, the specific intensity is always nonnegative. These corollaries ensure that Eqs. (44), (47), and (48) are physically meaningful.

7. Eqs. (44), (47), and (48) are easily generalized to the case of an incident field in the form of a superposition of several polychromatic parallel beams with quasi-monochromatic components and arbitrary propagation directions.

8. The quantity $i(\mathbf{r}, \mathbf{q})$ is nothing but a formal solution of the intermediate Eq. (46) and appears merely as a byproduct of the mathematical derivation of Eqs. (44) and (47) from the frequency-domain MMEs for a sparse random particulate medium. As such, it cannot be interpreted as describing the angular distribution of electromagnetic energy flow at the point $\mathbf{r}$. Fundamental physical significance can be ascribed only to the integral of $\mathbf{q}/(\mathbf{r}, \mathbf{q})$ over all directions $\mathbf{q}$, Eq. (44), rather than to the values of $i(\mathbf{r}, \mathbf{q})$ corresponding to individual directions. For example, adding to $i(\mathbf{r}, \mathbf{q})$ any function $f(\mathbf{r}, \mathbf{q})$ such that

$$\int_{4\pi} d\mathbf{q} \mathbf{q}/(\mathbf{r}, \mathbf{q}) = 0$$

yields another “specific intensity” causing the same $\langle S(\mathbf{r}, t) \rangle$, a simple example being any symmetric function such that $f(\mathbf{r}, -\mathbf{q}) = f(\mathbf{r}, \mathbf{q})$. We have seen before that even the Poynting vector cannot be legitimately claimed to specify the direction of time-averaged electromagnetic energy flow; obviously, there is even less rationale for attributing any “directional energy flow” content to the specific intensity.

9. If the particles forming the scattering medium are nonabsorbing, then it follows from the RTE that the time-averaged Poynting vector is divergence free:

$$\nabla \cdot \langle \mathbf{S}(\mathbf{r}, t) \rangle = 0. \tag{49}$$

This means that the time-averaged amount of electromagnetic energy entering a differential volume element per unit time is equal to the time-averaged amount of electromagnetic energy leaving the differential volume element per unit time. Since the RTE follows from the frequency-domain MMEs only upon making several well-defined assumptions, including the consideration of only the ladder diagrams, Eq. (49) shows that these assumptions are sufficiently consistent with each other in that the final result complies with the energy conservation law. Furthermore, it implies that the contribution of all the other types of diagram to the time-averaged Poynting vector must also be divergence free in the case of nonabsorbing particles.

10. Electromagnetic scattering can be caused not only by particles with distinct boundaries, but also by density and anisotropy fluctuations in rarified molecular media such as gases. Eqs. (44)–(48) remain valid in the case of scattering by a pure gaseous medium or a gaseous medium containing randomly distributed particles provided that all density/anisotropy fluctuations and particles are located in the far zones of each other.

11. It is straightforward to generalize the theory of electromagnetic scattering and the microphysical derivation of Eqs. (44)–(48) in order to account for absorption in the host medium \cite{190–192}.

12. The ladder approximation can be expected to work well in the near zone of a sparse particulate medium. In the far zone of the medium as a whole, the so-called maximally crossed (or cyclical) diagrams exemplified by Fig. 20e must be taken into account \cite{178,193} since they cause the effect of weak localization of electromagnetic waves in the backscattering direction \cite{127,139,193–200}.

5. Concluding remarks

Fig. 22 shows a diagram summarizing Section 4 and tracing the place of the microphysical theory of radiative transfer in particulate media within the broader context of classical Maxwell’s electromagnetics. Although I have been using the adjective “microphysical” to emphasize analytical back-traceability to the MMEs, it can also be said that this theory, as well as the theory of weak localization, is part of mesoscopic physics in that it deals with a size regime that is intermediate between the microscopic and macroscopic and is characteristic of a region where a large number of particles can interact in a correlated fashion. Direct computer solutions of the MMEs described in Refs. \cite{201,202} demonstrate indeed how the macroscopic regime of radiative transfer and weak localization emerges from the microscopic particle-level regime of Maxwell’s electromagnetics upon averaging over random realizations of a multi-particle group. Instructive
discussions of mesoscopic optical phenomena can be found in Refs. [199,200].

It is imperative to keep in mind that the formal applicability of the microphysical theory of radiative transfer rests on the specific approximations discussed in the preceding section, including the assumption that the scattering particles are located in each other’s far zones and are uncorrelated. The violation of this assumption in the case of densely packed particles can lead to significant errors in numerical predictions based on the RTT. Therefore, it is important to examine to what extent the RTT can be applied to densely packed particulate media. Initial results based on numerically exact solutions of the MMEs and controlled laboratory experiments have been reported in Refs. [202–205].

There is no doubt that the developments summarized in Section 4 amount to a profound paradigm shift. They reveal the inherently tensorial nature of electromagnetic energy transport in particulate media, clarify the statistical-optics content of the RTT and the physical nature of measurements with actual directional radiometers, and establish the microphysical disciplines of directional radiometry and radiative transfer in particulate media as valid and well-defined branches of physical optics. Of course, an interesting question is why these developments and their recognition by the scientific community have been so slow.

Part of the answer is provided by the following quote from Born and Wolf’s Principles of Optics [118]:
It seems to be a characteristic of the human mind that familiar concepts are abandoned only with the greatest reluctance, especially when a concrete picture of the phenomena has to be sacrificed.

Born and Wolf refer to the long struggle that Maxwell’s electromagnetic theory had to endure before it had become a generally accepted fundamental paradigm.\(^{10}\) However, the continued unwillingness to give up the intuitively appealing yet heuristic and even unphysical concepts discussed in Section 3 shows that the above quote has much broader relevance.

The existence of the microphysical theory of radiative transfer and directional radiometry in sparse particulate media does not necessarily imply that the corresponding phenomenological disciplines will fall out of circulation any time soon. In 1965, Rudolph Preiser predicted that even if a bridge connecting the fundamental physics and the island of the phenomenological RTT were built, one should not expect to see much traffic across this bridge [28]. To understand why this prediction has proved to be prophetetic, one can open the standard graduate-level textbook on radiative heat transfer published in 2013 [61]. The third edition of this popular text, advertised as “a comprehensive reference for scientists, engineers, and graduate students”, was “updated to include significant advances and the emergence of new research topics over the past decade”. However, even though various elements of the microphysical RTT have been known for half a century, they are nowhere mentioned. Furthermore, this textbook has only one chapter on the electromagnetic wave theory which, according to the Preface, “can (and will be) skipped by most instructors for a first course in radiative heat transfer”.

All in all, the resilience of the phenomenological paradigms of directional radiometry and radiative transfer has been quite enigmatic and represents an instructive proving ground for those applying the principles summarized in Thomas Kuhn’s Structure [27] to explain the direction and rate of scientific progress as well as illustrate concepts such as ”paradigm paralysis”. In fact, it might even be of special interest to a new generation of philosophers who go far beyond the scope of Ref. [27] in exploring the social character of scientific knowledge and scientific inquiry [206].

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References


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\(^{10}\) For example, Lord Kelvin, who died in 1907, had never accepted Maxwell’s theory of electromagnetism.