Credibility building activities in computational research include verification and validation, reproducibility and replication, and uncertainty quantification. Though orthogonal to each other, they are related. This paper presents validation and replication studies in electromagnetic excitations on nanoscale structures, where the quantity of interest is the wavelength at which resonance peaks occur. The study uses the open-source software PyGBe: a boundary element solver with treecode acceleration and GPU capability. We replicate a result by Rockstuhl et al. (2005, doi:10/dsxw9d) with a two-dimensional boundary element method on silicon carbide (SiC) particles, despite differences in our method. The second replication case from Ellis et al. (2016, doi:10/f83zcb) looks at aspect ratio effects on high-order modes of localized surface phonon-polariton nanostructures. The results partially replicate: the wavenumber position of some modes match, but for other modes they differ. With virtually no information about the original simulations, explaining the discrepancies is not possible. A comparison with experiments that measured polarized reflectance of SiC nano pillars provides a validation case. The wavenumber of the dominant mode and two more do match, but differences remain in other minor modes. Results in this paper were produced with strict reproducibility practices, and we share reproducibility packages for all, including input files, execution scripts, secondary data, post-processing code and plotting scripts, and the figures (deposited in Zenodo). In view of the many challenges faced, we propose that reproducible practices make replication and validation more feasible.
1. Introduction

Some fields of research, particularly solid mechanics and computational fluid dynamics, have a long tradition of community consensus building and established practices for verification and validation of computational models. Such practices are uncommon in other fields of science, especially if they have more recently become computationally intensive. Verification and validation also become increasingly difficult when the computational models arise from many levels of mathematical and physical modelling, representing a complex system. In recent years, science as a whole has come to be concerned with reproducibility and replication as a new front in the continuing campaign to build confidence on published findings. Together with formal processes of uncertainty quantification (UQ), we have now three complementary ‘axes’ for building trust in science.

The lengths to which research communities should go to conduct activities in verification and validation, reproducibility and replication, and UQ, are highly debated. Some journals require articles reporting on computational results to include proof of these activities, while most do not consider these aspects at all in their review criteria. In this paper, we tackle a sub-field of computational physics where tradition for these confidence-building activities is scant. The physical setting, excitation of resonance modes in nanostructures under an electromagnetic field, relies on multiple levels of modelling, while the experimental methods are complicated by the small length scales. We previously developed a computational model and software (called PyGBe) that has undergone code and solution verification, but a validation opportunity had remained elusive. Here, we present replication studies and a validation case based on published simulation and experimental results. Moreover, the studies in this paper were conducted under rigorous reproducibility practices, and all digital artefacts needed to reproduce every figure are shared in reproducibility packages available in a GitHub repository and archival services.

2. Background and methods

(a) Verification, validation, reproducibility and replication

Verification and validation of computational models—often abbreviated V&V and viewed in concert—have developed into a mature subject with more than two decades of organized efforts to standardize it, dedicated conferences, and a journal. The American Society of Mechanical Engineers (ASME), a standards-developing organization, formed its first Verification and Validation committee (known as V&V 10) in 2001, with the charter: ‘To develop standards for assessing the correctness and credibility of modelling and simulation in computational solid mechanics’. It approved its first document in 2006: The Guide for Verification and Validation in Computational Solid Mechanics (known as V&V 10-2006). The fact that this guide was 5 years in the making illustrates just how complex the subject matter, and building consensus about it, can be. Since that first effort, six additional standards sub-committees have tackled V&V in a variety of contexts. V&V 70 is the latest, focused on machine-learning models. The key principles laid out in the first V&V standard persevere through the many subsequent efforts that have operated to this day. They are:

— Verification must precede validation.
— Needs for validation experiments and associated accuracy requirements for computational model predictions are based on the intended use of the model.
— Validation of a complex system should be pursued in a hierarchical fashion from the component level to the system level.
— Validation is specific to a particular computational model for a particular intended use.
— Validation must assess the predictive capability of the model in the physical realm of interest, and it must address uncertainties that arise from both simulation results and experimental data.

The process of verification establishes that a computational model correctly describes the intended mathematical equations and their solutions. It encompasses both code correctness, and solution accuracy. Validation, on the other hand, seeks to determine to which measure a computational model represents the physical world. We like to say that ‘verification is solving the equations right, and validation is solving the right equations’ [1]. But in reality, the exercise can be much more complicated than this sounds. Computational models in most cases are built in a hierarchy of simplifications and approximations, and comparing with the physical world means conducting experiments, which themselves carry uncertainties.

As we will discuss in this paper, verification and validation in contexts that involve complex physics at less tractable scales (either very small, or very large), or where experimental methods are nascent, proceeds in a tangle of researcher judgements and path finding. In practice, validation activities reported in the scholarly literature often concentrate on using a stylized benchmark, and comparing experimental measurements with the results from computational models on that benchmark. Seldom do these activities address the key principles of pursuing validation in a hierarchical fashion from the component to the system level, and of assessing the predictive capability of the computational model accounting for various sources of uncertainties. Comprehensive validation studies are difficult, expensive and time consuming. Often, they are severely limited by practical constraints, and the conclusions equivocal. Yet the computational models still provide useful insights into the research or engineering question at hand, and we build trust on them little by little.

Verification and validation align on one axis of the multi-dimensional question of when are claims to knowledge arising from modelling and simulation justified, credible, true [2]. Two other axes of this question are: reproducibility and replication, and UQ. UQ uses statistical methods to give objective confidence levels for the results of simulations. Uncertainties typically stem from input data, modelling errors, genuine physical uncertainties, random processes, and so on. A scientific study may be reproducible, the simulations within it undergone V&V, yet the results are still uncertain. Building confidence in scientific findings obtained through computational modelling and simulation entails efforts in the three ‘axes of truth’ described here.

Reproducibility and replication (we could call it R&R) have preoccupied scientific communities more recently. Agreement on the terminology, to begin with, has been elusive [3]. The National Academies of Science, Engineering and Medicine (NASEM) released in May 2019 a consensus study report on Replicability and Reproducibility in Science [4] with definitions as follows. ‘Reproducibility is obtaining consistent results using the same input data, computational steps, methods, and code, and conditions of analysis. Replicability is obtaining consistent measurements or results, or drawing consistent conclusions using new data, methods, or conditions, in a study aimed at the same scientific question.’ According to these definitions, reproducibility demands full transparency of the computational workflow, which at the very least means open code and open data, where ‘open’ means shared at time of publication (or earlier) under a standard public license. This condition is infrequently satisfied. The NASEM report describes how a number of systematic efforts to reproduce computational results have failed in more than half of the attempts made, mainly due to inadequately specified or unavailable data, code and computational workflow [5–8]. Recommendation 4-1 of the NASEM report states that ‘to help ensure reproducibility of computational results, researchers should convey clear, specific, and complete information about any computational methods and data products that support their published results in order to enable other researchers to repeat the analysis, unless
such information is restricted by non-public data policies. That information should include the
data, study methods and computational environment’ [4].

Although it may seem evident that running an analysis with identical inputs would result
in identical outputs, this is sometimes not true. For example, the combination of floating-point
representation of numbers and parallel processing means that running the same software with
the same input data may give different numerical results. In some research settings, it may make
sense to relax the requirement of bitwise reproducibility and settle on reproducible results within
an accepted range of variation (or uncertainty). This can only be decided, however, after fully
understanding the numerical-analysis issues affecting the outcomes—and the risk associated with
an uncertainty range. Researchers using high-performance computing thus recognize that when
different runs with the same input data produce slightly different numeric outputs, each of these
results is equally credible, and the output must be understood as an approximation to the correct
value within a certain accepted uncertainty.

Beyond the particulars of high-performance or parallel computing, a number of factors can
contribute to the lack of reproducibility in research. In addition to lack of access to non-public
data and code, the NASEM report (of which Barba is a co-author) lists the following contributors
to lack of reproducibility:

— Inadequate record-keeping: the researchers did not properly document all relevant digital
artefacts and steps followed to obtain the results, the details of the computational
environment, software dependencies, and/or identifiers and metadata for data products.
— Lack of transparency: the researchers did not transparently report, using standard public
licenses, an archive with all relevant digital artefacts necessary to reproduce the results.
— Obsolescence of the digital artefacts: over time, the digital artefacts in the research
compendium are compromised because of technological breakdown and evolution or
lack of continued curation.
— Flawed attempts to reproduce others’ research: the researchers who attempted to
reproduce the work lacked expertise or failed to correctly follow the research protocols.
— Barriers in the culture of research: lack of resources and incentives to adopt reproducible
and transparent research across fields and researchers.

Improving computational reproducibility hinges on capturing and sharing information about the
computational environment and steps required to collect, process and analyse data. Scientific
workflows represent a complex flow of data products through various steps of collection,
transformation and analysis to produce an interpretable result. Capturing provenance of the
result is increasingly difficult to do using manual processes, and automation is key. With regards
to software management, version-control systems are used to automatically capture the history
of all changes made to the source code of a computer program. This creates a history of changes
and allows the developers to better understand the code and to identify possible problems or
errors. Recent technological advances in version control, virtualization, computational notebooks
and automatic provenance tracking have the potential to simplify reproducibility, and tools have
been developed that leverage these technologies. Still, many questions remain unanswered both
to understand the gaps left by existing tools and to develop principled approaches that fill those
gaps.

Replication of scientific findings is key for building trust in them. It is often difficult to attain,
for many reasons, not least because deciding when two scientific findings are consistent is tangled
in researcher judgements and inevitable constraints. The NASEM report lists the following factors
affecting the replicability of findings:

— the complexity of the system under study;
— understanding of the number and relations among variables within the system under
study;
— the ability to control the variables;
— levels of noise within the system (or signal to noise ratios);
— the mismatch of scale of the phenomena and the scale at which it can be measured;
— stability across time and space of the underlying principles;
— fidelity of the available measures to the underlying system under study (e.g. direct or indirect measurements); and
— prior probability (pre-experimental plausibility) of the scientific hypothesis.

Fields of study that have been at the centre of the ‘replication crisis’ commotion tend to be characterized by their complexity, intrinsic variability, or inability to control variables, e.g. psychology. But in many areas of modern technology, we face similar challenges to control variables or disentangle many interacting effects. In this paper, we tackle replication and validation of computational models in nanoscale physics, where certainly the systems are complex, variables difficult to control and signals subject to noise.

(b) Description of the PyGBe software

Our research group has been developing PyGBe—pronounced pigbeb—for several years. It was first written for biomolecular-electrostatics calculations using a continuum model of proteins in water or an ionic solvent. The computational model applies a boundary integral form of the governing Poisson–Boltzmann and Poisson equations, to obtain the electrostatic potential and its normal derivative on the molecular surface. In the implicit-solvent model, this information is used to compute the quantity of interest: solvation energy, which is diagnostic for questions of protein binding affinity, protein-surface interactions and others. Biomolecules are modelled as dielectric cavities inside an infinite continuum solvent, and the partial differential equations are solved via a boundary element method, leading to a dense linear system of equations. PyGBe uses a fast-summation algorithm via a Barnes-Hut treecode [9], and accelerates the computationally intensive parts of the code on Nvidia GPU hardware using CUDA kernels in the treecode, interfacing with PyCUDA [10]. Other portions of the code are written in C++, wrapped using swig, to extract more performance [11]. These features allow PyGBe to handle problems with half a million boundary elements or more.

In more recent work, we expanded the capabilities of PyGBe to applications of computational nanoplasmonics for biosensing [12]. Applying the long-wavelength limit, one can model the phenomenon of localized surface plasmon resonance (LSPR) via electrostatics, instead of the full Maxwell equations. This phenomenon is harnessed in nanosensors for detecting with high sensitivity the presence of biomolecules through shifts in resonance frequency. In LSPR, an electromagnetic wave excites the free electrons on the surface of a metallic nanoparticle. The name given to these electron-cloud vibrations is plasmons. In this setting, they resonate with the incoming field, and the energy is either absorbed by the nanoparticle or scattered in all directions, causing what’s called extinction. Since the resonance frequency depends on the dielectric environment, the change produced by a biomolecule approaching the metallic nanoparticle results in a frequency shift, and a means of detecting its presence. The electrostatic approximation allows using the methods implemented in PyGBe, after substantial code development to permit using complex-valued permittivities, and to incorporate an external electric field into the model. These changes included re-writing the Krylov iterative solver to work with complex numbers, and splitting the treecode evaluation into real and imaginary parts. New functions were added to read from configuration files the data describing the electric field, to compute the dipole moment, and to compute the final quantity of interest: extinction cross-section. We reported a major new release of the software in 2017 [13], and later presented the mathematical formulation for electromagnetic scattering in the long-wavelength setting and its associated continuum and discretized integral equations, and results including verification activities and sensitivity calculations on a biosensor model [12]. For verification, we conducted grid-convergence studies in two settings. In the first, we set up a computational model for a spherical silver nanoparticle in a constant electric field, leading to an analytical solution for the extinction cross-section. The second case does not have a
closed form solution: a spherical nanoparticle with a nearby protein (analyte), under an electric field. To estimate relative errors in the extinction cross-section, in this case, we made use of the method of Richardson extrapolation. These verification activities build our confidence on the computational model—moreover, our work was published following careful reproducibility practices, including the release of reproducibility packages for all results. Nevertheless, the gold standard of confidence in the predictive capability of the computational model comes from validation: our quest for an opportunity to conduct validation studies with PyGBe in these settings led us through the twisted path that we report in this paper.

(c) Physics context for this work

In recent years, polar dielectric crystals such as silicon carbide (SiC) became recognized as an alternative to plasmonic metals in many technologies, including biosensors. They manifest oscillations of lattice-bound charges, called surface phonon polaritons, in the mid- to long-wave infrared range with low optical losses. Nanostructures made of these materials offer sensing capabilities, described by their *figure of merits*, that are unattainable with plasmonic metals. The *figure of merits* of a nanoparticle is defined as the ratio between the sensitivity and the width of the resonance peak at mid-height [14]. In turn, sensitivity is the shift in the resonance position divided by the change in the refractive index: \[ S = \frac{\Delta \lambda}{\Delta n}. \] The dielectric function of polar dielectrics has a negative real part and a small imaginary part, in the mid- to long-infrared regime. This dielectric behaviour is similar to that observed in metals like silver in the blue part of the wavelength spectrum. In plasmonics, when illuminating a small particle made of metallic materials, we will observe that certain wavelengths excite a surface plasmon. The main difference with polar dielectrics like SiC is that for this material the frequency of the incoming light matches instead the resonance frequency of the Si and C sub-lattices [15,16]. This excitation leads to a strong extinction cross-section at the resonance wavelength, and an enhanced near-field amplitude. These behaviours can be modelled with the same approaches used for localized surface plasmon resonance, and when the wavelength is much larger than the size of the nanoparticle, we can again apply the electrostatic approach implemented in PyGBe [12,13].

When studying both surface phonon polaritons and surface plasmon resonances, one can analyse the spectrums by measuring different quantities. We can measure scattering cross-section, absorption, or extinction cross-section (scattering plus absorption), as well as reflection. Since the *quantity of interest* is the wavelength (frequency) at which the resonance modes happen, these different approaches are comparable in some cases, e.g. whether we measure reflection or extinction cross-section, the wavelength (frequency) at which the peaks happen remains the same. Throughout this work, we will concentrate on studying the wavelength (frequency) at which the resonance modes occur, and we aim to replicate results from Rockstuhl *et al.* [16] and from Ellis *et al.* [17], and to validate our software against experimental results from Ellis *et al.* [17].

3. Results

All the results reported in this paper were obtained using the PyGBe software [13], with the version at commit 34eddfe (https://github.com/pygbe/pygbe/tree/e1f3650b0c99cab99dce5372200d3a1534eddfe) in the history. The software GitHub repository contains a Dockerfile to create the container image where we ran the simulations. The manuscript GitHub repository is separate from the software repository, and can be found at https://github.com/barbagroup/pygbe_validation_paper. It contains the reproducibility packages for all results, as described in §3c and elsewhere in the paper. We used a laboratory workstation for all simulations, built from parts. Hardware specifications are as follows:

— CPU: Intel Core i7-5930K Haswell-E 6-Core 3.5 GHz LGA 2011-v3
— RAM: G.SKILL Ripjaws 4 series 32 GB (4 \( \times \) 8GB)
— GPU: Nvidia Tesla K40c (with 12 GB memory)
Solver parameters: We used a GMRES exit tolerance of $1 \times 10^{-6}$ in the iterative solver for all our simulations. Details on the treecode and integration parameters (and others) can be found within input files included in the manuscript GitHub repository as part of the reproducibility packages (repro-packs).

Run times: We report detailed time logs together with the data of our simulations, as part of the repro-packs to accompany this paper. Here is a brief report. For the result of the replication of fig. 14 of Rockstuhl et al. §a, the total wall-clock time for producing each curve was approximately 11.6 h, which is the result of computing the extinction cross-section for 175 wavelength cases, giving $\approx 4$ min per run. In the case of the validation and replication of fig. 2a of Ellis et al. §b, the total wall-clock time for producing the curve was approximately 2.3 h, which is the result of computing the extinction cross-section for 208 wavelength cases, giving $\approx 40$ s per run.

(a) Replication of results from Rockstuhl et al. [16]

The work of Rockstuhl et al. [16] studies the phonon-polariton response of SiC nanoparticles using a two-dimensional (2D) boundary element method, previously developed in their group [18]. They analysed ‘cylindrical particles’ (where they extend in the third dimension to infinity) made of 6H-SiC, with varying cross-sections. We decided to attempt to replicate one of the results presented on fig. 14 of their paper: the scattering cross-section of a SiC rectangular cylinder for three different aspect ratios. To be well within our quasi-static approach ($\lambda > d$ where $d$ is the characteristic dimension of the geometry), we chose the case with $a = 672$ nm, $b = 328$ nm.

(i) Differences in method, mesh and dielectric data

Method: The main difference between the original simulations and ours is that they solve a 2D problem with the full Maxwell equations, while we solve a 3D problem with the electrostatic approximation. We lack any information about their code implementations, discretization schemes or solver. We computed extinction cross-section (scattering plus absorption) while Rockstuhl et al. present only scattering cross-section. Absorption dominates in intensity over scattering in the quasi-static regime, so these results are not generally comparable. When the resonance peaks are sharp and narrow, however, the wavelengths at which they occur in the scattering and absorption spectra are nearly the same. For example, Wiley et al. show simulation results for a silver nanocube where the main peaks in the spectra are close [19]. Since silicon carbide has more pronounced peaks than silver, the wavelength at which they occur in the extinction and scattering spectra are more closely aligned.

Mesh: Rockstuhl et al. did not provide any details regarding the discretization of the geometries or parameters involved in the simulations. We performed a grid-independence study as a form of solution verification, and to ensure that we were minimizing errors due to discretization. Rockstuhl et al. used a 2D geometry (infinite third dimension), while we treat the geometry in its full 3D representation.

Dielectric data: The study uses 6H-SiC as material, and the authors obtained their data from a source that we were not able to replicate. Instead, we used experimental data for 4H-SiC that was provided to us via private communications from the authors of Ellis et al. [17].

(ii) Grid-independence study

We performed a grid-independence study on a SiC cube of side $L = 535$ nm submerged in air, under a constant electric field aligned with the z-axis (a similar set-up to the square cylinder on fig. 18 of Rockstuhl et al. [16]). Due to the nature of the geometry and its sharp edges, it was challenging to see proper convergence. We have completed grid-convergence analysis for this type of physics in a previous work, but using spherical geometries [12]. With that experience, and given the difficulty caused by sharp edges, we are content with studying grid-independence here instead. Figure 1 shows the grid-independence study, using meshes with 15 552 triangles (density
Figure 1. Grid-independence study for a SiC cube of side $L = 535$ nm submerged in air under a constant electric field in the $z$-direction. The curves represent the extinction cross-section divided by $L^2$ as a function of wavelength divided by $L$, for mesh sizes 19 K = 19 200 triangles and 15 K = 15 552 triangles. The label 'diff' refers to the difference between the results of the two meshes. (Online version in colour.)

9.05 $\times$ $10^{-5}$ per Å squared) to 19 200 triangles (density 1.11 $\times$ $10^{-4}$ per Å squared): the computed results show no discernible difference.

It is worth noting that the extinction cross-section curve in figure 1 has extra peaks compared to the results of fig. 18 of Rockstuhl et al. due to the three-dimensional effects captured in our simulation and the sharp edges, the latter effect also being mentioned by Rockstuhl et al. The 3D effects will be addressed in the following results, where we attempt a replication of one of the results on fig. 14 of Rockstuhl et al.

(iii) Replication of fig. 14 (case a1) of Rockstuhl et al. [16]

We chose to replicate a result of Rocksuthl et al. presented in fig. 14: the case where $a = 672$ nm and $b = 328$ nm, since these dimensions are well within the limits of the quasi-static approximation used in PyGBe. Rockstuhl et al. present the normalized scattering cross-section of a SiC rectangular cylinder, and they perform simulations for two different set-ups, sketched in figure 2. In their fig. 14 (left), they have the wavevector (illumination) along the long side of the geometry, which means that the electric field is parallel to the short side of the rectangle, like in fig. 2b. Following a similar analysis, in fig. 14 (right) of Rockstuhl et al. they have the wavevector (illumination) along the short side of the geometry, which means that the electric field is parallel to the long side of the rectangle, like in fig. 2a.
(b) Replication of results from Ellis et al. [17], and validation

The work of Ellis et al. [17] studies the aspect-ratio evolution of high-order modes in localized surface phonon-polariton nanostructures. They study the excitation of multipolar localized surface phonon polaritons (SPhP) resonances, by measuring and computing polarized reflectance...
Figure 4. Effect of uniformity of the mesh and roundness of the edges on the extinction cross-section of a rectangular prism of SiC of dimensions $a = 672$ nm, $b = 328$ nm and $y = 2688$ nm, submerged in air and under a constant electric field parallel to the z-axis. The labels are: Trimesh, for a non-uniform mesh generated using Trimesh; Uniform, for a uniform mesh generated using Python scripts; and Uni + round, for a uniform mesh generated using Python scripts with round edges obtained using Trimesh. (Online version in colour.)

Figure 5. Replication of the results in fig. 14 of Rockstuhl et al. [16]. Extinction cross-section of a rectangular prism of SiC of dimensions $a = 672$ nm, $b = 328$ nm and $y = 2688$ nm, submerged in air and under a constant electric field parallel to the z-axis (PyGBe). The digitized curve from Rockstuhl et al. corresponds to scattering (Rockstuhl). (Online version in colour.)

on 4H-SiC pillars of fixed height ($H = 950$ nm), fixed width ($W = 400$ nm) and varied length ($L = 400–4800$ nm). These pillars are patterned on a square grid with a pitch $P = L + 500$ nm to reduce coupling. In both their simulations and experiments, they measured polarized reflectance with the incident polarization oriented parallel or perpendicular to the long axis of the pillars.

Our first aim was to replicate the computational result presented in figure S4 of their supplementary material, corresponding to the black curve on their plot. In this figure, they show simulation results for the resonance spectral position of the lower-frequency mode when having parallel polarization ($E_{\parallel}100$), with an incidence angle of 22 degrees. We attempted to replicate this result since the gap between the pillars is 5000 nm (10 times larger than in the other cases), diminishing the effects of coupling, which makes it a better candidate to replicate using PyGBe. In our calculations, the set-up consists of a single pillar, with no substrate. Our second attempted replication is for the results presented in fig. 2a, corresponding to reflectance measurements...
across wavenumber for pillars with aspect ratio $AR = 4$, angle of incidence 22 degrees, and incoming polarization parallel to the length of the pillars. For this case, the authors also reported experimental results that we will use for validation of our solver.

(i) Differences in method, mesh and dielectric data

Method: Ellis et al. ran experiments using reflectance spectroscopy and they computed the solution of Maxwell’s equations via the RF package of the finite-element solver in the commercial software COMSOL. In their simulations, they used one pillar over a portion of substrate, with periodic boundary conditions to represent an array of pillars and their interactions. We use the boundary element method in the quasi-static approximation, which is suitable in this case since the pillar’s size is small compared to the wavelengths involved in the simulations: in the range 10000–12500 nm. We measured extinction cross-section, which will express as peaks instead of dips (shown in the reflection plots of Ellis et al.). The intensity of the peaks is not comparable, but we are looking to match the wavenumber (quantity of interest) at which these events happen.

Mesh: For the case with aspect ratio $AR = 4$, we have a non-uniform triangular surface mesh ($N = 4398$) provided to us by the authors of Ellis et al., which we used for validation and replication of fig. 2a of their paper. However, for the replication of figure S4, we needed the remaining aspect ratio meshes. We generated uniform meshes using our Python script, and we determined the density of these meshes by comparing the extinction cross-section for the case with $AR = 4$ of our mesh and the one provided by Ellis et al. We noted that using around double the number of elements ($N = 8564$) than in the original mesh and rounding the edges, the relative errors for the extinction cross-section were, on average, smaller than 3%, and the variation on the wavelength peak position was smaller than $1 \text{ cm}^{-1}$. This analysis led us to a density of $\approx 1.7 \times 10^{-5}$ triangles per Å squared to create the meshes for the aspect ratio variation study.

Dielectric data: The dielectric data for the simulations was given to us by the authors of the paper via a private communication (CTE & JGT, 2019), and corresponds to experimental values of the dielectric.

(ii) Replication of figure S4 in the supplementary materials of Ellis et al. [17]

To be able to replicate the result of figure S4 of Ellis et al., we need to identify the lower-frequency mode for each of the aspect ratios in our computations. For each aspect ratio (AR) value from 1 to 7, we computed the extinction cross-section $C_{ext}$ across the wavenumbers in the range 800–1000 cm$^{-1}$. We identified the lower-frequency mode ($E_{\parallel 100}$ for Ellis et al.) that is not a longitudinal mode. The longitudinal modes appear only when we have an incidence angle that is off-normal, since they are associated with the height of the pillar. Figure 6 shows the results of the extinction cross-section of a SiC pillar of fixed height ($H = 950 \text{ nm}$), fixed width ($W = 400 \text{ nm}$) and varied length ($L = 400–2800 \text{ nm}$), for normal and 22-degrees angle of incidence. The simulations were performed for the long-edge orientation, meaning that the electric field is aligned with the length of the pillar when having normal incidence as shown on figure 7. From these results, we selected the lowest mode that is not a longitudinal one and extracted its wavelength (table 1) to replicate figure 8. This figure shows the results from Ellis et al. (digitized by hand using the WebPlotDigitizer) and the results obtained with PyGBe, and table 2 shows that the percentage error is below 2% for all cases. (Note that Ellis et al. changed the angle of incidence of the illuminating vector while in PyGBe we rotated the geometry instead.)

(iii) Validation of PyGBe against experimental results in fig. 2a of Ellis et al. and replication of the corresponding computations

The results of fig. 2a in Ellis et al. were obtained on a pillar of aspect ratio $AR = 4$. For this case, we have the original mesh, provided to us by the authors. We also know that our computation for the mode $E_{\parallel 100}$ compares well with theirs (percentage error 0.16%), therefore, we attempted to validate our simulations with their experimental results (red curve on their paper), as well as to
Figure 6. Extinction cross-section across wavenumbers for SiC pillars of varying aspect ratios, \((H = 950 \text{ nm}, W = 400 \text{ nm}, L = 400–2800 \text{ nm}, \text{AR} = 1–7)\), with both normal incidence and a 22-degree incidence. (Online version in colour.)
**Figure 7.** Diagram showing the angles of incidence in our simulation set-ups to comply with the configuration in the case from Ellis *et al.* (Online version in colour.)

**Figure 8.** Replication of figure S4 in the supplementary materials of Ellis *et al.* [17]. Wavenumber at which the $E_{100}^\parallel$ mode happens for different aspect ratios. (Online version in colour.)

**Figure 9.** PyGBe versus the experiments presented in fig. 2a of Ellis *et al.* [17] (we obtained the data digitizing by hand from their figure using WebPlotDigitizer). (Online version in colour.)
Table 1. Wavelength at which peaks happen for different aspect ratios, for runs where the electric field is parallel to the length (L) of the pillar. We have normal incidence and 22-degree incidence. The wavelengths in bold correspond to the lowest mode that is not a longitudinal one.

| AR  | % error | 917.73  | 934.092 | 949.604 | 957.325 |
|-----|---------|---------|---------|---------|---------|
| 22° | 983.926 | 917.73  | 935.052 | 949.604 | 957.325 |
| 22° | 903.233 | 926.395 | 944.762 | 958.242 |
| 3   | 888.793 | 922.552 | 931.223 | 948.613 | 958.242 |
| 22° | 888.793 | 899.418 | 922.552 | 931.223 | 958.242 |
| 4   | 876.186 | 929.32  | 946.639 | 958.242 |
| 22° | 876.186 | 901.281 | 921.618 | 929.32  | 945.745 | 958.242 |
| 5   | 865.576 | 926.395 | 945.745 | 958.242 |
| 22° | 865.576 | 901.281 | 921.618 | 958.242 |
| 6   | 856.904 | 914.793 | 923.489 | 929.32  | 946.639 | 958.242 |
| 22° | 856.904 | 901.281 | 920.6   | 958.242 |
| 7   | 850.134 | 910.963 | 921.618 | 928.372 | 946.639 | 958.242 |
| 22° | 850.134 | 901.281 | 910.963 | 920.6   | 958.242 |

Table 2. Percentage error for different aspect ratios.

| AR  | % error |
|-----|---------|
| 1   | 0.95    |
| 2   | 0.67    |
| 3   | 0.35    |
| 4   | 0.16    |
| 5   | 0.72    |
| 6   | 1.20    |
| 7   | 1.59    |

replicate their simulations (green curve on their paper). Figure 2r of Ellis et al. presents measured and simulated reflectances of SiC pillar arrays with a 500-nm gap. All their measurements and simulations were performed with 22° off-normal angle of incidence and incoming polarization parallel to the elongated side of the pillar.

Using PyGBe, we computed the extinction cross-section of an isolated SiC pillar (AR = 4) with no substrate, submerged in air under a constant electric field in the z-direction, and rotated the orientation of the pillar to match the angle of incidence used by Ellis et al. (figure 7). In figure 9, we present a comparison of our simulations and the experimental results of Ellis et al. A difference in the wavelengths of the peaks is noticeable. This may be attributed to the fact that in their experiments the separation between pillars is 500 nm, which implies there are coupling effects that in our simulations are not considered.

First-order correction. Since our simulations do not take into account coupling effects in an array of prisms, we cannot strictly match the conditions to validate our solver. However, from figure S4
in the supplementary materials of Ellis et al., we know that coupling effects affect the $E_{100}^{||}$ mode by a shift of 12.17 cm$^{-1}$. Therefore, as a first-order correction we can subtract this amount from our simulations to account for coupling effects. In figure 10, we present the result after applying this correction for coupling effects. It is worth mentioning that the far-left (837 cm$^{-1}$) and far-right (964 cm$^{-1}$) peaks on the results of Ellis et al. are peaks associated with zone-folded LO (longitudinal) phonons of 4H-SiC, an effect they say is beyond the scope of their analysis [17]. They concentrate their analysis on the peaks that occur between 864 and 961 cm$^{-1}$. If we use the same approximation and compare the results with the simulations of Ellis et al. on fig. 2 (green curve on their paper), we obtain figure 11.

(c) Reproducibility and data management

Barba [20] describes the elements of reproducible computational research that we have adopted in our practice. A key element is professional software management and engineering, and the central
technology solution is version control. Preserving a complete history of changes is the only way to manage complex software projects that can support reproducibility. Moreover, all our research software is developed in the open, and shared under permissive public licenses, such as BSD-3 or MIT License. These open licenses permit all uses, as well as derivative works, only subject to attribution. Another key element of transparent and reproducible computational modelling is automation. Automating every step means turning protocols into code. For example, simulations are launched with parameters fed from a configuration file rather than an interactive prompt, and all figures and plots are produced by writing scripts, rather than pointing-and-clicking on a graphical interface. The goal is to create complete ‘recipes’ in code, which can be run, versioned and shared.

Our signature reproducibility practice is to organize and share in an archival-quality repository (providing a global identifier) all digital artefacts associated with the results in the paper. This includes input files, configuration files, post-processing scripts, files specifying the build process and containerization (e.g. Docker files), and even cloud computing machine definitions, if applicable [21]. We call these openly shared file sets reproducibility packages, or repro-packs for short, and we have been doing this for many years and improving the process iteratively. The basic steps were already contained in the ‘Reproducibility PI Manifesto’ of 2012 [22], where Barba pledged to always share a manuscript’s data, plotting scripts and figures under CC-BY (Creative Commons Attribution license). Notably, this makes the figures re-usable by readers, without requiring them to ask permission from the publisher, even if there was a transfer of copyright of the paper (the figures are included in the paper under the conditions of the public license). We later extended the practice to include all other digital artefacts associated with the results, beyond secondary data and figures. As in our previous papers, readers can reproduce all the figures in this paper using the repro-packs shared in the manuscript GitHub repository, and archived separately in Zenodo. They include Jupyter notebooks with all the plotting code, and also the manually digitized data from the figures in the source articles for our replication cases. Barba & Thiruvathukal [23], explain that it is not enough to provide these materials in a GitHub repository (the owner of a repository is always able to delete it), and one should deposit the reproducibility packages in an archival service providing a global identifier and permanent link (like a digital object identifier, DOI).

The following items of archived digital artefacts accompany this paper:

— The software repository for PyGBe is at https://github.com/pygbe/pygbe.
— The repository for this paper is at https://github.com/barbagroup/pygbe_validation_paper, which also includes the manuscript source files in LaTeX.
— The problem datasets for replication of Rockstuhl et al. [16] are in the manuscript repository, but also archived in Zenodo at 10.5281/zenodo.3962534 [24].
— Execution files for all runs are archived in Zenodo at 10.5281/zenodo.3962576 [25].
— The problem datasets for validation and replication of results from Ellis et al. [17] are archived in Zenodo at 10.5281/zenodo.3962584 [26].
— The file sets for reproducing the figures for the replication of results from Rockstuhl et al. [16] are archived in Zenodo at 10.5281/zenodo.3962791 [27].
— The file sets for reproducing the figures for the validation and replication of results from Ellis et al. [17] are archived in Zenodo at 10.5281/zenodo.3962797 [28].

4. Discussion

We have presented several results that replicate previous published findings in the general area of nanostructure responses to electromagnetic waves. Our field of interest is computational nanoplasmonics for applications in biosensors, and in a previous publication we developed the mathematical formulation and reported both solution verification activities, and an application demo with our software PyGBe, extended to treat complex dielectrics and imposed electric fields [12]. The search for a physical context and published results that would allow us to undertake
validation studies with PyGBe is what led us to this work. Even if we finally do have validation and replication cases, neatly presented here, the path to obtain these results was nonlinear, iterative and arduous.

In the first case, we sought to replicate a result from Rockstuhl et al. [16], where they computed the scattering cross-section as a function of wavelength for a silicon carbide rectangular nanostructure. They present their results as a plot (fig. 14 in their paper), and report the numeric value of the resonance wavelengths in the text. Lacking access to the secondary data behind the plots—computed from two-dimensional simulations with a boundary element solver—we were forced to manually digitize the values from the figure. Our results are presented in figure 5, together with the curve we obtain from digitizing the source image. We were successful at replicating the strong peaks reported by Rockstuhl et al. at wavelengths 10.42 µm and 10.7 µm, when the electric field $E$ is parallel to the short side of the rectangle, and 10.42 µm and 10.82 µm when $E$ is parallel to the long side. Our results contain extra (small) peaks that are not present in the work of Rockstuhl et al. The first one, located between the main two peaks, we attribute to the effect of sharp edges (figure 4), as when we introduce a level of roundness, it diminishes. The second extra peak is the far right one, and we believe this peak is a consequence of the 3D nature of our geometry; as observed in figure 3, this peak intensity decreases as the third dimension of the geometry lengths. The quantity of interest in these findings is the wavelength of the resonance peaks, and our results do indeed match the findings.

The second replication case is from a paper by Ellis et al. [17] studying the effect of aspect ratio on the excitation of high-order modes in localized surface phonon-polariton nanostructures (figure S4 of their supplementary material and fig. 2 of their paper). Figure 8 shows the results for this replication, where the relative errors between our computations and theirs is always smaller than 2%. The smallest error is for the case with AR = 4 and we proceeded to study the results on fig. 2 of their work, corresponding to this same aspect ratio. Their results in fig. 2a include both experiments and simulations with the commercial software COMSOL, so we sought to both validate PyGBe using their experimental results, and replicate their computational findings. Again, we lack access to the data behind the plots, and we had to digitize the curves by hand. The quantity plotted in the original figures is reflectance as a function of wavenumber, whereas we compute the extinction—on the figures, they show inverted peaks, where we show positive peaks. We can compare the results, nevertheless, because the quantity of interest is the wavenumber position of the peaks. Figure 9 shows the results of our simulations using PyGBe on an isolated pillar, compared with the experimental results of Ellis et al. on an array of pillars. The results with PyGBe do not account for the effect of coupling among the pillars, which explains the noticeable discrepancy on the wavenumbers at which the peaks occur. We proposed a correction, based on the results reported on Figure S4 of Ellis et al. for the AR = 4 case, where the shift on the wavenumber due to coupling is 12.17 cm$^{-1}$ (difference between black and red curves for AR = 4 on figure S4 of the supplementary materials). We subtracted this value to that obtained with our simulations and we show the comparison of our corrected results against those of Ellis et al.: the experimental data on figure 10, and the computational data on figure 11. When comparing with their experiments, after the correction was applied, we observe a good match of the wavenumber for the lower (and stronger) mode, as well as a good match for the third and fourth peaks. The wavenumber of the second peak, related to a longitudinal excitation (mode $L_{001}$ in Ellis et al.), presents a discrepancy that we believe is related to the fact that our pillar does not have a substrate underneath. The remaining (fifth) peak also presents a discrepancy, but in this case, we cannot identify the reason. We did not analyse the peaks out of the range 864–961 cm$^{-1}$, since Ellis et al. describe these peaks to be associated with zone-folded LO (longitudinal) phonons of 4H-SiC, and outside the scope of their study. After considering all these details, we can say that we have validated our solver PyGBe against the experimental results of fig. 2a, as well as replicated their computational results.

Throughout all these activities aiming to replicate previous results and validate our computational model, we faced multiple challenges, starting with the complexity of the system under study. In both our source papers, we have systems that we were unable to fully model
using PyGBe. For the case of Rockstuhl et al. even though they used a boundary element method, it was a 2D model instead of 3D like the one implemented in PyGBe. The computational work presented in Ellis et al. used a volumetric formulation (finite-element method), whereas PyGBe implements a boundary integral one. Both works computed (or measured) quantities different from the extinction cross-section (computed in PyGBe), but given the relationship between their quantities (scattering cross-section and reflection) and ours (extinction cross-section), via the wavelength (wavenumber) at which resonance happens, this was not a problem. To produce the end results for comparing against these works, however, we went through an exhaustive process of modelling, making assumptions and even corrections. We were lacking any information regarding the solvers, discretization, meshes, etc., in the original papers. In the case of Ellis et al., we benefited from multiple communications with the authors, who made available the geometry and mesh of the pillar, as well as the dielectric data used as input for the simulations. Even though we appreciate this helpful interaction, replication studies should ideally not depend on communications with the original authors. In both cases, the secondary data presented in their results were not publicly available and we relied on a manually digitized version, obtained from the plots. This work is tedious and introduces uncertainties that would be avoided by releasing the secondary data.

Validation has been a hard goal to achieve for our solver since the field does not have benchmarks that are meant to be used for this purpose. Multiple times, we encountered experimental results that we aspired to use for validation, but due to the insufficient reporting of the experimental conditions, the set-up and the geometries involved, we have been unable to validate until this moment. Even though we were not modelling the same experimental set-up, the communications with the authors of Ellis et al. made the validation possible.

After all the challenges faced to achieve code validation and replication of results, we can say that the process was far from linear, and the complexity involved increases if the work to be replicated or used for validation was not conducted and published using reproducible practices, and made open at the time of publication. We conclude that reproducible practices are needed not only for the work to be reproduced but also replicated and even used for validation studies, if applicable.

Data accessibility. https://github.com/barbagroup/pygbe_validation_paper https://doi.org/10.5281/zenodo.3962534 https://doi.org/10.5281/zenodo.3962576 https://doi.org/10.5281/zenodo.3962584 https://doi.org/10.5281/zenodo.3962791 https://doi.org/10.5281/zenodo.3962797.

Authors’ contributions. N.C. and L.B. contributed to research design. N.C. carried out computations. N.C. and L.B. wrote the paper.

Competing interests. We declare we have no competing interests.

Funding. This research is supported by the National Science Foundation under Award no. 1747669 of the Computer and Information Science and Engineering (CISE) Directorate.

Acknowledgements. We thank Dr Chase T. Ellis and Dr Joseph G. Tischler for the fruitful discussions regarding their results [17], which led to the validation of PyGBe, and for providing a mesh file and dielectric data that made the modelling closer to their original set-up. We also thank Dr Christopher D. Cooper for many valuable discussions that helped us overcome some of the challenges faced throughout this work. After we posted the preprint on arXiv (just after submission to the journal), we received communication from Prof. Dr Carsten Rockstuhl that helped us clarify some points in our paper, which we appreciate.

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