Quantum ESPRESSO toward the exascale

Paolo Giannozzi,1,2,3 Oscar Baseglio,4 Pietro Bonfà,5,6 Davide Brunato,4 Roberto Car,7 Ivan Carnimeo,4 Carlo Cavazzoni,8,3 Stefano de Gironcoli,4,6 Pietro Delugas,4,3 Fabrizio Ferrari Ruffino,2 Andrea Ferretti,6 Nicola Marzari,9,3 Iurii Timrov,9 Andrea Urru,4 and Stefano Baroni4,2,3

1) Dipartimento di Scienze Matematiche, Informatiche e Fisiche, Università di Udine, via delle Scienze 206, I-33100 Udine, Italy, European Union
2) CNR-IOM, Istituto dell’Officina dei Materiali, SISSA, I-34136 Trieste, Italy, European Union
3) Quantum ESPRESSO Foundation, Cambridge Road Ind Estate, Milton, Cambridge, CB24 6AZ, United Kingdom
4) SISSA – Scuola Internazionale Superiore di Studi Avanzati, via Bonomea 265, I-34136, Trieste, Italy, European Union
5) Dipartimento di Scienze Matematiche, Fisiche e Informatiche, Università di Parma, Parco Area delle Scienze 7/A, I-43124 Parma, Italy, European Union
6) Centro S3, CNR-Istituto Nanoscienze, via Campi 213/A, I-41125 Modena, Italy, European Union
7) Department of Chemistry, Princeton University, Princeton, NJ 08544, USA
8) CINECA - Via Magnanelli 6/3, I 40033 Casalecchio di Reno, Bologna, Italy, European Union
9) Theory and Simulation of Materials (THEOS), and National Centre for Computational Design and Discovery of Novel Materials (MARVEL), École Polytechnique Fédérale de Lausanne (EPFL), CH-1015 Lausanne, Switzerland

(Dated: 23 April 2021)

QUANTUM ESPRESSO is an open-source distribution of computer codes for quantum-mechanical materials modeling, based on density-functional theory, pseudopotentials, and plane waves, and renowned for its performance on a wide range of hardware architectures, from laptops to massively parallel computers, as well as for the breadth of its applications. In this paper we present a motivation and brief review of the ongoing effort to port QUANTUM ESPRESSO onto heterogeneous architectures based on hardware accelerators, which will overcome the energy constraints that are currently hindering the way towards exascale computing.

I. INTRODUCTION

The goal of this manuscript is to describe recent and ongoing work on the QUANTUM ESPRESSO software distribution for first-principle atomistic simulations. We focus in particular on the challenges posed by the new heterogeneous architectures, based on multi-core chips enhanced with multiple hardware “accelerators”, coupling exceptional performances to an acceptable energy consumption. The large-scale adoption of these emerging architectures across different classes of computing systems will bring a paradigm shift into scientific computing, similar to what vector machines caused in the 80’s, parallel machines in the 90’s, massively parallel machines more recently.

This paper is organized as follows.

In Sec. II, we briefly describe the history and the current status of QUANTUM ESPRESSO. We give a short overview of its features and capabilities, by mostly referring to the relevant literature.

In Sec. III we describe the challenges posed and opportunities offered by heterogeneous architectures. The opportunity is to reach what is dubbed “exascale computing”: an unprecedented amount of computer power, opening new perspectives to computer simulations. The challenges, for scientific software in general and for QUANTUM ESPRESSO in particular, are unprecedented as well. The amount of needed changes is much larger, and the effects of changes much deeper, than in previous transitions to vector, parallel, massively parallel architectures. Moreover, several competing architectures are hitting the marketplace, each one coming with a different software stack and tools.

Sec. III A describes ongoing work towards performance portability, that is: the ability to obtain comparably high performances on different computer architectures, minimizing the need for maintaining hardware-specific code versions. The work of Sec. III A, that focuses on low-level libraries, would however be futile without more work at a higher level, aimed towards future maintainability of a large and complex scientific software project.

Sec. III B deals with recent and ongoing work aiming at a sustainable development model, that is: restructuring codes in a way that makes them easier to maintain, to extend, and especially to port to other architectures in a foreseeable future.

Sec. III C describes the current status, capabilities, and some benchmarks for QUANTUM ESPRESSO on NVIDIA graphics processing units (GPUs), one of the leading candidate architecture for the future “exascale computer”. The benchmarks are designed for relatively small machines and do not aim at showing performances on large-sized systems. They aim instead at pointing out bottlenecks, inefficiencies, and the minimum size of calculations that saturate the computational power of a GPU.

Finally, Sec. IV contains our conclusions: a brief analysis of the achieved results and an outlook on forthcoming actions.

II. QUANTUM ESPRESSO AT THE TURN OF THE TWENTIES

The QUANTUM ESPRESSO project was started in 2002, with the merger of three packages for density-functional the-
ory (DFT) simulations using plane waves and (ultrasoft) pseudopotentials, which had been under development since the mid-80’s:

- **PWscf**: code pw.x for self-consistent field (SCF) solution of Kohn-Sham equations and structural optimization, code ph.x for lattice-dynamical calculations using linear response, plus many other utility codes;
- **CP**: a code performing first-principle molecular dynamics (MD) simulations of the Car-Parrinello type, specialized to large supercells;
- **FPMD**: a code similar to CP, but with a different and partially overlapping set of functionalities.

Innovation in quantum-mechanical materials modeling has always been one of the main concerns and distinctive features of this project. In particular, the linear-response code ph.x contained in PWscf was, to the best of the authors’ knowledge, the first implementation of density-functional perturbation theory (DFPT). The elimination of virtual orbitals from linear-response and excited-state calculations, pioneered in this implementation, has remained one of the distinctive features of QUANTUM ESPRESSO, later to be adopted by several other codes. On the same line, CP and FPMD were both offsprings of the first implementation of Car-Parrinello MD, now merged into a single code, cp.x.

Since the beginning, QUANTUM ESPRESSO was released under an open-source license (GNU GPL) and was conceived as a “distribution”: an integrated suite of packages, following loose programming guidelines, rather than a monolithic application. The rationale for these choices lies in the philosophy of fostering collaboration and sharing of code among scientists, creating a community of users and developers, while pursuing innovation in methods and algorithms. For more on such aspect, we refer to Refs. 8 and 9.

The QUANTUM ESPRESSO distribution has much evolved over the years. On the one hand, more sophisticated theoretical methods and algorithmic advances have been implemented, in particular:

- projector-augmented waves (PAW);
- non-collinear and spin-orbit coupling (SOC) calculations;
- Hubbard-corrected functionals;
- nonlocal functionals and other semi-empirical and less-empirical corrections for van der Waals bonded systems;
- hybrid functionals, also exploiting orbital localization.

On the other hand, QUANTUM ESPRESSO has extended its original scope with additional packages for more property calculations. We mention in particular:

- activation energies and reaction barriers using the nudged-elastic-band (NEB) method;
- superconducting transition temperatures, electrical resistivity, and other effects related to electron-phonon interactions;
- phonon linewidths, thermal conductivity, and anharmonic effects;
- nuclear magnetic resonance (NMR) chemical shifts and electron paramagnetic resonance (EPR) g-tensors;
- transport in nano-wires in the ballistic regime;
- near-edge X-ray absorption spectra;
- implicit solvation and electrochemical models;
- optical and electron energy loss spectra using time-dependent density-functional perturbation theory.

On the purely computational side, optimization for modern and massively parallel high-performance computing (HPC) architectures has been pursued, using multiple parallelization levels involving both message-passing interface (MPI) and multi-threading (OpenMP).

All these advances, and related references, are documented in Refs. 8 and 9. In particular, Ref. 9 also documents a preliminary version of package HP, performing linear-response calculations of Hubbard parameters, that has been since released.

In addition to being directly used for implemented property calculations, QUANTUM ESPRESSO is used in more creative ways as well:

- as a “quantum engine” for more sophisticated calculations, such as advanced MD calculations implemented in i-PI, genetic and evolutionary algorithms, high-throughput calculations with AiIDA;
- in modified or patched versions, for specific purposes, as in “embedded” eQE.

Finally, QUANTUM ESPRESSO is used to produce DFT data for further processing by other codes. We mention in particular codes performing Quantum Monte Carlo calculations (QMCPack) and many-body perturbation theory (MBPT) (Yambo, BerkeleyGW, WEST, Sternheimer-Gw): determining maximally localized Wannier functions (Wannier90 and electron-phonon interactions (EPs)); and codes performing different types of data analysis (e.g., topological analysis with critic2) or computing transport properties (e.g., Wann, KGEC).

Stable versions of the source code—the latest is v.6.5 at the time this article was written—as well as development and experimental versions can be downloaded from the URL:

http://www.quantum-espresso.org/download.

### III. CHALLENGES AND OPPORTUNITIES OF NEW HETEROGENEOUS ARCHITECTURES

The push of quantum mechanical materials simulation software towards ever increasing levels of complexity, accuracy, and performance has been so far assisted by a constant downsizing of micro-processing units, allowing for a steady increase of the compute capacity of general-purpose architectures at constant power. This process is now hitting its ultimate physical limits and is just about to come to an end. To reverse this state of affairs, major and disruptive changes in hardware architecture are to occur. The constraints set by power consumption can only be met by heterogeneous architectures, with specialized cores (“accelerators”) that maximize efficiency for a small set of instructions: e.g., graphics processing units (GPUs), PEZY chips, tensor processors, neuromorphic chips, etc. A quick look at the first places of the Top500 supercomputers list clearly shows that heterogeneous architectures have become the *de facto* standard for new gen-
On the single node scale, the ratio between the computational power provided by accelerators and traditional CPU units is found to grow at each procurement cycle over the last 10 years. Most of the computational power of future “exascale” machines (that is: capable of 1 exaflop, or $10^{18}$ floating-point operations per second) will come from accelerators.

The ensuing architectural complexity will set demanding requirements in terms of data movement, heterogeneous memory management, fault tolerance, which will all require a major, possibly joint, re-design of circuits and algorithms and the adoption of different programming paradigms. In particular, extremely parallel applications will require rapid and substantial architectural shifts, including, for example, the handling of intra-node data movement between disjoint memory spaces and the explicit treatment of deeper memory hierarchies.

Porting community codes to novel hardware architectures has always required extensive re-coding in the past. This cannot be sustained any longer in view of the considerable complexity reached by QUANTUM ESPRESSO and similar community codes (several hundred thousands code lines each) and the forthcoming diversity, heterogeneity, and rapid evolution of the hardware architectures. The solution we have identified is to refactor QUANTUM ESPRESSO into multiple layers, resulting from the assembly of weakly coupled components (modules and libraries), to be maintained and enhanced independently from each other, shared among different codes, and designed to be as architecture-agnostic as possible. A bird’s eye view of the code will reveal four main such layers (see Fig. 1):

- The top layer is a collection of property calculators to compute materials properties and to simulate processes, which is the ultimate goal of molecular and materials simulations. These property calculators may share global variables among themselves and with the quantum-engine modules (see below). This layer could be partially and progressively made code-agnostic, while it should remain as architecture-agnostic as possible.

- The core layer contains quantum-engine modules that solve the one-particle Schrödinger equation and perform associated tasks, such as Hamiltonian builds (i.e. the application of the one-particle Hamiltonian to molecular orbitals and Bloch states), or other tasks related to density-functional, linear-response, and many-body perturbation theories. This layer is and will likely continue to be code-specific, but should stay as architecture-agnostic as possible. Ideally, the same quantum-engine modules could be shared among different codes performing similar tasks, e.g. standard SCF computations and geometry optimizations, ab initio MD, and linear-response/MBPT calculations.

- A collection of domain-specific mathematical libraries, to perform general-purpose tasks, such as 3D Fourier analysis, linear algebra, using both iterative and factorization algorithms, non-linear optimization, etc. Such mathematical libraries should be easily shared among different codes of a same class (e.g. adopting the same quantum-mechanical representation / basis set) and stay largely architecture-agnostic as above. While these libraries may themselves contain modules, in order to ensure easy portability across different codes, the data exchange between them and the calling code will only occur via standard public application programming interfaces (APIs), which will make minimal use of complex data structures and derived data types.

- Finally, a collection of low-level system libraries, which will abstract the most recurrent architecture-specific constructs, such as data offload to/from specialized hardware, memory synchronization, and the like. These libraries will interface directly with the underlying hardware architectures and should be fine-tuned on each of them for optimal performance. They should be architecture-specific, and as much code-agnostic as possible, without interfering with the other layers.

This strategy is meant to accomplish what has come to be dubbed separation of concerns: ideally, method developers in science departments and research labs should be concerned...
with property calculators, disregarding architectural details, whereas scientists and research engineers in IT departments and HPC centers should focus on low-level mathematical and system libraries. The two categories of developers should work shoulder-to-shoulder to develop and maintain the quantum engine. Separation of concerns is the overarching guideline for the action of the EU MAX Centre of Excellence (CoE) for HPC Applications, whose mission is to foster the porting of important community codes for quantum materials modeling to heterogeneous architectures.

The implementation of the above strategy is bound to entail an extensive refactoring of community codes, which in turn will require a longer time than allowed by the pressing advance of heterogeneous architectures. For this reason, the course of action undertaken by the QUANTUM ESPRESSO developers follows two converging tracks. On the one hand, an accelerated version of pw.x working on NVIDIA GPUs is already available, and more components—initially cp.x, followed by ph.x and other linear-response codes—are being ported. On the other hand, the refactoring of the whole distribution into a collection of stand-alone modules and libraries is steadily progressing. The final goal to which both tracks will eventually converge is the identification and gradual encapsulation of architecture-specific computational kernels into low-level libraries, so as to make the migration to multiple architectures as seamless as possible. We aim in particular at using the forthcoming OpenMP-5 standard, that in addition to architecture-agnostic APIs (already in OpenMP 4.5) provides deep memory copies and a simple framework for exploiting unified shared memory when available.

In the following we discuss in some detail the different logical layers that constitute the structure of the QUANTUM ESPRESSO distribution, as shown in Fig. 1, starting from the bottom layer. Low-level system libraries are discussed in Sec. III A while the remaining layers are discussed in Sec. III B.

A. Performance portability

Performance portability across current and future heterogeneous architectures is one of the grand challenges in the design of HPC applications. General-purpose frameworks have been proposed, but none of them has reached maturity and widespread adoption. In addition, Fortran support is still very limited or missing entirely. In this context, the MAX CoE is promoting and coordinating a collective effort involving the developers of various materials modeling applications. Taking on this challenge with a domain-specific approach has the advantage of providing abstraction and encapsulation of a limited number of functionalities that constitute the building blocks of the most common operations performed on the accelerators in this field. This will allow us to prepare low-level architecture-specific implementations of a limited number of kernels that have been already characterized and isolated, thus keeping the source code of the various scientific applications untouched and reducing code branches when new systems will appear on the market.

Such an effort is still in the early stages, but is under active development and is progressively entering the GPU port of QUANTUM ESPRESSO through the so-called DevXlib library. This library started off as a common initiative shared among MAX CoE codes (notably QUANTUM ESPRESSO and Yambo), aimed at hiding CUDA Fortran extensions (see Sec.III C) in the main source base. Being used by different codes, the library has been rationalized and further abstracted, thus becoming a performance portability tool aimed at supporting multiple back-ends (support to OpenACC and OpenMP-5 foreseen, direct extension to CUDA C possible). The main features included in the library by design are the following:

- performance portability for Fortran codes;
- deal with multiple hardware and software stacks, programming models and missing standards;
- wrap/encapsulate device specific code;
- focused on limiting code disruption (to foster community support).

It is important to note that part of the library design includes the definition of which device-related abstract concepts need to be exposed to the scientific developers. To give an example, memory copy and synchronization to/from host/device memory are abstract operations that the developers of property calculators or of the quantum engine itself may need to control directly. Therefore, DevXlib exposes such control in the form of library APIs that are agnostic of the specific device backend.

In practice, DevXlib provides the user with (i) interfaces to memory handling operations including creation and locking of memory buffers (device_memcpy and device_buffers); (ii) interfaces to basic and dense-matrix linear algebra routines, similarly to BLAS and Lapack (device_linalg); (iii) interfaces to more domain-specific operations (device_auxfuncs); (iv) device-oriented data structure compatible with Fortran usage. In particular, memory handling allows the user to copy memory host-to-host, device-to-device, and also across memories, host-to-device and vice-versa, thereby dealing also with memory off-load and synchronization. Importantly, both synchronous and asynchronous copies can be performed with explicit control. Moreover, the explicit handling of memory buffers is meant to ease or avoid the procedure of allocation and deallocation of auxiliary workspace.

Besides the interface to linear algebra optimized libraries, such as cuBLAS and cuSOLVER in the case of NVIDIA GPUs, DevXlib also provides interfaces to selected (more specialized) kernels that appear often in plane waves electronic structure codes (such as scalar products with G-vector remapping or specific matrix or vector updates, to name a few). While not covering all possible kernels of the same complexity, this is quite effective in helping to keep the code sources as clean as possible. Importantly, some efforts have also been directed to devise Fortran-compatible data structures to handle the extra complexity of data stored on host and/or accelerator memories. In principle, these data structures need to be able to hide hardware complexity (e.g. being vendor agnostic for what concerns accelerator memory), to
allow for seamless memory synchronization and transfer, and
to be equipped with a set of methods to implement the most
common data operations. Software engineering and exper-
imentation of these data structures is currently ongoing.

In order to make the code more flexible and portable, some
basic functionalities are accessed via common interfaces pro-
vided by a low-level library *UtilXlib*, including utilities for
MPI and OpenMP parallelization, timing, error and memory
handling. This library has been extended to include execution
and data synchronization points for communications involv-
ing memory spaces located on the GPUs.

B. Towards a sustainable development, maintenance, and
porting model

*QUANTUM ESPRESSO* has grown in size during the
years, including as of v.6.5 almost 600,000 lines of Fortran
code, 60,000 lines of C, python, shell scripts, plus a large
amount of tests, examples, and documentation. While not es-
pecially large with respect to scientific software projects in
other fields, *QUANTUM ESPRESSO* is sufficiently bulky to
make its maintenance and extension a challenge that cannot
be sustained without resorting to modern software engineer-
ing techniques. Much work, described in part in Sec. 3 of
Ref. 9, has been done in the past along the following direc-
tions:

- provide a large set of automated tests to ensure the va-
  lidity and portability of the results under different build-
  ing and execution patterns;
- extend the interoperability with other codes via struc-
tured I/O, using an extended markup language (XML)
  with an industry-grade schema description for small
  human-readable, data files, and optionally a hierarchi-
cal format (HDF5) for large, binary, data sets;
- collect large parts of the code base into modules and li-
  braries, in order to enhance its readability, ease of main-
  tenance, and portability.

The work along the latter direction has been extended with the
creation of distinct code layers: *(i)* Libraries and *(ii)* Modules.
*Libraries* have a well-encapsulated inner data structure and
exchange data with the rest of the code only through prede-
finite APIs. As a consequence, libraries can be developed, dis-
tributed, and compiled independently of each other and then
linked to the main code. *Modules* are reusable blocks of code
whose functionalities are accessed via well-defined APIs as
well but, mainly for reasons of computational efficiency, do
not stick to such a clear-cut data encapsulation scheme as li-
braries and share a significant portion of their data structure
with the rest of the code. For this reason, modules must be
compiled together with the main code and are mainly intended
for usage inside the *QUANTUM ESPRESSO* suite or other
codes sharing the same global data structure.

1. Domain-specific mathematical libraries

Apart from improving the maintainability of the whole dis-
tribution, packaging the code base into libraries also has the
advantage of making distributed development and mainte-
nance easier and of providing a wider community with effec-
tive tools for developing electronic-structure software ready to
use in modern HPC infrastructures. For this reason we aim at
avoiding the usage of structured data type as arguments of the
APIs as much as possible and at exposing the interfaces using
included files rather than Fortran modules.

Currently, three major packages have been extracted from
*QUANTUM ESPRESSO* and are ready to be distributed as
stand-alone libraries, namely:

- *LAXlib*, performing parallel dense-matrix operations,
  including basic linear algebra and diagonalization;
- *FFTXlib*, for parallel distributed three-dimensional fast
  Fourier transforms;
- *KS_Solvers*, a collection of iterative diagonalization
  algorithms to solve the Kohn-Sham equations.

A large part of the computations of a typical electronic-
structure calculation is performed inside these libraries. The
usage of machine-optimized mathematical libraries and the
inclusion of further optimizations, either architecture-agnostic
or architecture-specific, in these libraries will automatically
profit to all codes and computations. It is at this level that
the separation of concerns is most fruitful in terms of per-
formance portability. While the original code targeted MPI
and OpenMP parallelization on many CPUs, the extension to
different programming paradigms for heterogeneous archite-
tures has much progressed since, also thanks to contributions
from IT experts.

*LAXlib* and *FFTXlib* libraries, with their completely en-
capsulated inner data structures, can be easily used by third
parties. Their interfaces are transparent to the specific under-
lying architecture.

The iterative diagonalization algorithms collected inside the
*KS_Solvers* are disentangled from the specific Hamiltonian
builder, which is called by the library as an external routine;
the definition of wavefunctions and their scalar products ins-
ide the Hamiltonian builder must be compatible with the one
used inside *KS_Solvers*. For some of the algorithms, a Re-
verse Communication Interface (RCI) is also available, allow-
ing one to directly pass the $H |\psi\rangle$ vectors to the library, leav-
ing to the programmer the task of computing and converting
them to the format expected by the RCI.

The goals of the activities described here largely overlap
with those of the Electronic Structure Library (ESL) project at
CECAM. The possibility of decoupling the *KS_Solvers*,
*LAXlib* and *FFTXlib* libraries from their native codes was
first demonstrated during a Workshop organized in 2017
within the ESL initiative. Moreover, both *KS_Solvers* and
*LAXlib* may use another library maintained by ESL, ELPA (in-
cluded in ESL) for dense-matrix diagonalization. *QUANTUM
ESPRESSO* may also use the *libxc* ESL library computing
exchange-correlation functionals.
2. Quantum-engine modules

For other parts of the code, data encapsulation is difficult to achieve or even unfeasible, or may introduce inefficiencies. For those cases, it was chosen to refactor the code into general modules, still using the global data structure of the suite. These modules are currently meant to be used inside the distribution, but they are designed to be easily accessible for the development of further applications built upon the QUANTUM ESPRESSO code base. Notable examples are Modules and LR_Modules.

Modules is a collection of Fortran modules and subroutines that implement various operations needed to solve self-consistently and iteratively the Kohn-Sham equations of DFT. In particular, Modules contains the following functionalities: (i) definition of global variables and universal constants, (ii) reading of input parameters and of pseudopotentials, (iii) definitions of Bravais and reciprocal lattices, (iv) symmetry analysis and symmetrization operations, (v) calculation of the exchange-correlation potential, (vi) generation of plane waves and of their mapping to FFT grids, (vii) generation of k points, (viii) calculation of pseudopotential terms. Historically, Modules exists since the very first version of QUANTUM ESPRESSO, but it has continuously evolved in order to adapt to novel utilities and packages of the suite.

LR_Modules is a much more recent part of QUANTUM ESPRESSO, which appeared about five years ago and evolved significantly since that time. The reason for the creation of LR_Modules was to unify, harmonize, generalize, and refactor the functionalities that are common to all linear-response and MBPT codes of the suite. LR_Modules contains the following functionalities: (i) definition of global data structures for linear response, (ii) calculators of linear-response quantities (such as e.g. response density and potentials), (ii) iterative solvers (e.g. Lanczos recursive algorithms), (iii) response exchange-correlation kernel calculators, (iv) symmetrization routines, (v) projectors on the empty-states manifold, to name a few. The functionalities of LR_Modules are used in the following packages:

- PHonon for calculation of lattice vibrational modes (phonons). Born effective charges, dielectric tensor, and other vibrational properties;
- TDDFPT for calculation of optical absorption spectra of molecular systems; collective excitations in solids such as plasmons and magnons;
- EPW for calculation of electron-phonon coupling, transport, and superconducting properties of materials;
- HP for the first-principles calculation of Hubbard parameters of the Hubbard-corrected DFT.

The generalized and unified subroutines from LR_Modules have been refactored in such a way that they can be easily and straightforwardly employed in any other future linear-response or MBPT code of QUANTUM ESPRESSO or even in third-party codes. They can now be used generically to build perturbations, apply them to the occupied ground-state Kohn-Sham wave functions and compute the related self-consistent first-order response properties either by solving the Sternheimer equations or by solving the Liouville quantum equations using the Lanczos recursion method.

3. Interoperability

Exportable output in QUANTUM ESPRESSO is based on the adoption of standard data formats: XML and HDF5. These two formats have the advantage of providing information about the hierarchy and the types of the data that may be automatically processed by external applications. The support of these features in modern scripting languages like python makes them convenient for the development of postprocessing and analysis tools. For HDF5 the description of the hierarchy and of data types is contained in the file; for XML files we provide it under the form of XSD schemas. In order to streamline the reading of XML files in python using the specifications of the schemas we have also released a python package, xmlschema, that converts the ASCII content of the XML file into a corresponding python dictionary whose structure and data types follow the indications given by the schema. The python dictionary can then be used directly by the reading application or saved as JSON or YAML files.

The coherence between the released schemas for XML and the effective output of the applications is guaranteed by a python set of tools that produce Fortran bindings for reading and writing XML data, starting from the XML schema. The tools generate the writing routines from the format specification and keep them automatically aligned with the schema. These tools have also been released as a separate package.

The qeschema package provides the APIs specific for reading and writing the XML, HDF5 and unified pseudopotential format (UPF) files used by the applications in the suite. This package also provides converters to map some structured data as e.g. crystal structures to and from other format commonly used in visualizers, structure databases or atomistic simulation utilities as ASE or pymatgen or HDF5 charge densities which may be exported to other formats, for instance the XSF format of crystal.

The postQE package provides python extensions and APIs to use the postprocessing utilities of the PP package inside python applications. Many PP components are compiled as python extensions using F2PY. For those applications that may have a large computational load and memory footprint, postQE provides instead tools to extract data from the output files of the Fortran executables.

C. Evolution of the GPU-enabled version

QUANTUM ESPRESSO introduced support for accelerated systems as early as 2013, starting from v. 5.0, in the form of custom plugins for some of the codes of the suite. The initial approach, based on CUDA C and ad hoc libraries for linear algebra, proved to be successful in boosting the code performance, but hardly sustainable from the maintenance and development points of view, mainly due to the substantial amount of “boilerplate” (replicated) code used to interface Fortran subroutines with CUDA C.
In light of this limitation, a new port, dubbed QE-GPU, has been recently rewritten from scratch, starting from the case study presented by Romero et al.\textsuperscript{65} who ported v. 6.1 of \textit{pw.x} to NVIDIA GPUs.\textsuperscript{66} In Ref. 65, the authors detail a new strategy based on CUDA Fortran—the Fortran analogue of CUDA C—and demonstrate 2× to 3× speedup consistently achieved on a variety of platforms and using different benchmarks. The new GPU-enabled releases of \textsc{Quantum Espresso} extend this work, but adopting a few design solutions to streamline future development and porting to other heterogeneous architectures, as detailed below.

While still being developed with CUDA Fortran, the last release of QE-GPU is almost entirely accelerated through a directive-based approach, using the so-called \textit{cuF} kernel compiler directive, that generates parallel architecture-specific code for loop-based structures. In some cases this choice may limit the code performance, but it brings a number of positive consequences. Firstly, it allows one to validate GPU code on the CPU and, in principle, to retain a single source code for both CPU and GPU implementations. Second, this design choice enables the straightforward adoption of other directive-based programming models, like OpenACC or OpenMP, if required. As a consequence, even if the current implementation only runs on NVIDIA GPUs, special care has been paid to design the software in a way that minimizes the future effort needed to port the code to other accelerators that can be exploited through a directive-based programming model. In particular, attention has been paid not to introduce CUDA Fortran kernels in the high-level structure of the code, except for trivial data transformations, and to keep them confined to the domain-specific and system libraries described in Secs. III A and IIIB.

A further advantage of CUDA Fortran is explicit memory management. \textsc{Quantum Espresso} organizes related data into modules and derived types. Allocation and synchronization of these kinds of data structures is a straightforward and concise operation in CUDA Fortran, while OpenACC and OpenMP support automatic synchronization of these data types only in the most recent releases. In addition, by retaining full control on memory allocations and synchronizations between the RAM and GPU’s global memory, we provide a clear view of all intra-node memory operations to the programmer, thus facilitating future porting activities.

As detailed in Sec. III B, a few specific components of \textsc{Quantum Espresso} have been extracted from the main code trunk, made independent from it and released as packaged libraries. Some of them have been targeted for GPU acceleration, namely the parallel dense eigenvalue solver (LAX11b), and the parallel distributed FFT (FFTXX11b). In this case we abandoned directive based acceleration in favor of architecture specific APIs and libraries. Indeed, the last release of these libraries relies on cuBLAS for linear algebra kernels, cuFFT for 1D and 2D Fourier transforms, and cuSOLVER for solving real and complex generalized eigenvalue problems.\textsuperscript{67}

As of version 6.5, the following set of functionalities benefit from GPU acceleration and operate on data residing on the GPU:

- electronic self-consistency for both magnetic and spin-polarized systems;
- iterative solution of the Kohn-Sham Hamiltonian using either the Conjugate Gradient or the Davidson method;
- calculation of atomic forces;
- calculation of exact exchange terms for hybrid functionals.

The acceleration is obtained by exploiting numerous kernels that have been ported to GPU: local-potential, pseudopotential, and reciprocal space sampling is reported close to the chemical formula. For each simulation the sum of the time taken by the initialization, the iterations during self consistency and the estimation of atomic forces is compared.
tial, kinetic-energy contributions to the Hamiltonian, preconditioner evaluation for iterative solvers, application of the overlap operator, wavefunction initialization, Hubbard component contribution to the effective potential, charge density generation. For all the aforementioned operations, where applicable, both the case of real-valued wavefunctions ($k = 0$ only sampling, useful to reduce the memory footprint and to speed up simulations) and the case with spinor wavefunctions (non-collinear magnetism) have been ported to GPU.

Currently, only pw.x can benefit from GPU acceleration, but other codes are being ported and will be available in future releases. The GPU-enabled version of pw.x is fully compatible with its CPU counterpart, provides the same features, undergoes the same regression testing suite, and converges to equivalent results within a given convergence criterion.

The speedup provided by the GPU implementation depends drastically both on the hardware and on the details of the input data. The extreme scale performance has been already detailed elsewhere, thus here we focus on smaller problems, consisting of tens of atoms and hundreds of electrons. This will allow us to identify the lower limit for the input cases that can benefit from GPU acceleration.

The qeinputgenerator was used to prepare a set of benchmarks. This tool automatically generates input files for pw.x providing a few options to customize the accuracy of the simulation and using pseudopotentials from two sets, either standard solid-state pseudopotentials (SSSP) Efficiency or SSSP Precision. These in turn include norm conserving 72–74 ultrasoft, 75 and PAW 76,77 pseudopotentials thus covering a significant fraction of pw.x functionalities. Sixteen structures having unit cell volume between 250 Å³ and 1000 Å³ were randomly selected from the Crystallography Open Database (COD). Structures with fractional occupations and rare earth elements were discarded. All input and output data are available on the Materials Cloud archive.

In Figure 2 we compare the best time-to-solution obtained with a single 18-core Intel(R) Xeon(R) E5-2697 v4 @ 2.30GHz (BDW) CPU and the same hardware accompanied by one NVIDIA’s Tesla V100 GPU card. The ratio between the theoretical peak performance of these two units is roughly 1 to 10, but effective GPU acceleration can only be achieved with extremely data parallel workloads and the speedup provided by the graphic card can even become lower than 1 when this condition is not met. This limit is investigated with the present benchmark using a rather standard balance between GPU and CPU computational power for a HPC node.

The CPU results have been collected with QUANTUM ESPRESSO v. 6.5 compiled with Intel’s 2019 suite, Intel’s MPI implementation and Intel’s Math Kernel Library (MKL), while QE-GPU v. 6.5 was built with the PGI 19.10 Fortran compiler and linked to Intel’s 2019 MKL. For the set of inputs detailed above, the pure MPI parallelization is the best strategy for the CPU version of the code, therefore we performed all simulations for the CPU version disabling OpenMP. On the other hand, the GPU code requires OpenMP parallelism since the number of MPI processes is limited by the number of GPU cards installed on the system; each MPI process should be assigned just one accelerator. For both the CPU and the GPU versions, only parallelism over k-points (-NPOLL option) has been used: a reasonable choice, given the relatively small dimension of the dense eigenvalue problem to be solved during iterative diagonalization.

The two sets of results are numerically equivalent: the largest discrepancy between the total energies computed by the CPU and the GPU versions is $2 \times 10^{-8}$ Ry, while the largest difference in the total force is $3 \times 10^{-5}$ Ry/Bohr.

In Fig. 2 we report the wall time required to reach convergence (or 80 SCF cycles when convergence was not achieved) for the CPU and GPU versions. Only the best time to solution as a function of k-point parallelism is reported. The total execution time has also contributions, not reported because much smaller, from the initialization step and by the final calculation of atomic forces. For the smallest test case, a 98-atom organic compound with 246 electrons that converges in less than 1 minute on our hardware, the GPU speedup is just 1.4x. This is indeed the current limiting size for taking advantage of GPU acceleration. On the other hand, as the problem size grows, especially as a function of the basis set dimension, the speedup can exceed 3x.

IV. OUTLOOK AND CONCLUSIONS

The work on the QUANTUM ESPRESSO distribution has since many years followed some well-established lines of action: (i) implementing new numerical algorithms, theoretical methods, property calculations; (ii) improving interoperability with external codes; (iii) making the codes more modular and easier to work with, more portable across architectures, without sacrificing performances. The arrival of heterogeneous, accelerated architectures has made the latter direction more urgent and wider in scope. It is no longer sufficient to isolate low-level computational kernels into simple mathematical libraries: performance portability must be ensured at the level of so-called “domain-specific” libraries. Without effective actions in the direction (iii), it will become difficult to implement, and even more difficult to maintain, further property calculations, new theories, and new algorithms.

The work described in Sec III C is the starting point for the effective implementation of the plan described in Sec. III. The next significant step is the merger of the main QUANTUM ESPRESSO distribution with the CUDA Fortran version for NVIDIA GPUs. This will eliminate the constraint of keeping the latter version akin to the main distribution in order to simplify the “alignment” process as the main distribution evolves.

In parallel, more work is ongoing to achieve performance portability, as described in Sec. III A. In particular, support for OpenMP-5 is being introduced into the domain-specific mathematical libraries of Sec. III B and the low-level system libraries of Sec. III A. In this respect we are working with Intel software engineers to guarantee complete compatibility for their future Ponte Vecchio GPU architecture, and QUANTUM ESPRESSO will be ready to run on these cards when they will appear on the market, towards the end of 2021. Porting to AMD accelerated architectures is also ongoing exploiting the public OpenACC stack. Work on ARM vectorized (SVE
instruction set) architectures (EP188 and AFX6489 processors) is also ongoing and on track to release an optimized QUANTUM ESPRESSO version for these architectures. All these porting efforts would be hardly feasible without the contribution of IT experts and HPC centers, in the spirit of the separation of concerns. This solution does not target the maximum level of architecture-specific optimizations, but leaves the possibility open to achieve them, once the hardware is finally installed in major HPC centers, using specially-tailored patched versions.

The benchmarks presented in Sec. III C are realistic but relatively small-size use cases. The extension to large-scale calculations requires further work to identify and remove memory and computation bottlenecks. In particular, the amount of global memory per card is a relevant parameter that substantially impacts the performance of the accelerated version of the code. Some of the largest benchmarks in our set do saturate the memory available on a single card, depending upon the parallel options used to distribute plane-wave components. Similar issues will certainly show up also in extreme scale benchmarks. For single-point self-consistent or molecular-dynamics calculations, the most critical bottlenecks towards scaling to the whole exascale system will be the amount of memory in cards, due to the superlinear complexity of plane-wave electronic structure calculations (the amount of memory and computation increases more than quadratically with the size of the system). It is foreseen that the best “exasflop” performances will be actually achievable for selected cases, such as “high-throughput” and “ensemble” large-scale calculations, that can be split into many smaller ones. The efficiency of each computation will thus be critical for fully exploiting the capability of the HPC systems.

Data availability statement. The data that support the findings of this study are openly available in the Materials Cloud Archive at https://doi.org/10.24435/materialscloud:2020.0021/v1.84

ACKNOWLEDGMENTS

This work was partially funded by the EU through the MAX Centre of Excellence for HPC applications (Project No. 824143). Financial and administrative support from the QUANTUM ESPRESSO Foundation90 is also gratefully acknowledged. N.M. and I.T. acknowledge support from the Swiss National Science Foundation (SNSF), through grant 200021-179138, and its National Centre of Competence in Research (NCCR) MARVEL. Valuable technical support from NVIDIA is gratefully acknowledged. QUANTUM ESPRESSO has received contributions from many people in addition to the authors of this article. In particular, we would like to thank the following colleagues who have contributed to different stages of the design and implementation of the QE-GPU code: Fabio Affinito, Anoop Chandran, Brandon Cook, Massimiliano Fatica, Ivan Girotto, Thorsten Kurth, Miloš Marić, Everett Philips, Josh Romero, and Filippo Spiga.

1Stefano Baroni, Paolo Giannozzi, and Andrea Testa. Green’s Function Approach to Linear Response in Solids. Phys. Rev. Lett. 58:1861–1864, May 1987.
2P. Giannozzi, S. de Gironcoli, P. Pavone, and S. Baroni. Ab initio calculation of phonon dispersions in semiconductors. Phys. Rev. B, 43:7231, 1991.
3Stefano Baroni, Stefano de Gironcoli, Andrea Dal Corso, and Paolo Giannozzi. Phonons and related crystal properties from density-functional perturbation theory. Rev. Mod. Phys., 73:515–562, Jul 2001.
4X. Gonze. Abiadiatic density-functional perturbation theory. Phys. Rev. A, 52:1096–1114, Aug 1995.
5X. Gonze. Erratum: Abiadiatic density-functional perturbation theory. Phys. Rev. A, 54:4591–4591, Nov 1996.
6R. Car and M. Parrinello. Unified approach for molecular dynamics and density-functional theory. Phys. Rev. Lett., 55(22):2471–2474, Nov 1985.
7At the time, PaNeT had already been released under the GPL.
8Paolo Giannozzi, Stefano Baroni, Nicola Bonini, Matteo Calandra, Roberto Car, Carlo Cavazzoni, Davide Ceresoli, Guido L. Chiarotti, Matteo Coccioni, Ismaila Dabo, Andrea Dal Corso, Stefano de Gironcoli, Stefano Fabris, Guido Fratesi, Ralph Gebauer, Uwe Gerstmann, Christos Gougoussis, Anton Kokalj, Michele Lazzeri, Layla Martin-Samos, Nicola Marzari, Francesco Mauri, Riccardo Mazzarello, Stefano Paolini, Alfredo Pasquarello, Lorenzo Paulatto, Carlo Sanzacco, Sandro Scandolo, Gabriele Scandolo, Ari P. Seitsonen, Alexander Smogunov, Paolo Umari, and Rezende M Wentzcovitch. QUANTUM ESPRESSO: a modular and open-source software project for quantum simulations of materials. J. Phys.: Condens. Matter, 21(39):395502, 2009.
9Paolo Giannozzi, Oliviero Andreussi, Thomas Blumme, Oana Bnau, Marco Buongiorno Nardelli, Matteo Calandra, Roberto Car, Carlo Cavazzoni, Davide Ceresoli, Matteo Coccioni, Nicola Colonna, Ivan Carneiro, Andrea Dal Corso, Stefano de Gironcoli, Pietro Delugas, Robert A. DiStasio Jr., Andrea Ferretti, Andrea Floris, Guido Fratesi, Giorgia Fungai, Ralph Gebauer, Uwe Gerstmann, Feliciano Giustino, Tommaso Gorni, J. Jia, M. Kawamura, H.-K. Ko, Anton Kokalj, Emine Küçükbenli, Michele Lazzeri, Margherita Marsili, Nicola Marzari, Francesco Mauri, N. L. Nguyen, H.-V. Nguyen, Albert Otero-de-la Roza, Lorenzo Paulatto, Samuel Poncé, Dario Rocca, Riccardo Sabatini, Biswajit Santra, Martin Schlüpf, Ari Seitsonen, Alexander Smogunov, Iuriy Timrov, Timo Thonhauser, Paolo Umari, Nathalie Vast, Xifang Wu, and Stefano Baroni. Advanced capabilities for materials modelling with QUANTUM ESPRESSO. J. Phys.: Condens. Matter, 29:465901, 2017.
10Iuriy Timrov, Nicola Marzari, and Matteo Coccioni. Hubbard parameters from density-functional perturbation theory. Phys. Rev. B, 98:085127, 2018.
11Venkat Kapil, Mariana Rossi, Ondrej Marsalek, Riccardo Petraglia, Yair Litman, Thomas Spura, Bingqing Cheng, Alice Cuzzocrea, Robert H. Meilner, David M. Wilkins, Benjamin A. Helfrecht, Przemyslaw Juda, Sebastien P. Bienvenue, Wei Fang, Jan Kessler, Igor Poltavsky, Steven Vandervande, Jelle Wieme, Clemence Corninboeuf, Thomas D. Kühne, David E. Manolopoulos, Thomas E. Markland, Jeremy O. Richardson, Alexandre Tkatchenko, Gareth A. Tribello, Veronique Van Speybroeck, and Michele Ceriotti. i-PI 2.0: A universal force engine for advanced molecular simulations. Computer Physics Communications, 236:214 – 223, 2019.
12Giovanni Pizzi, Andrea Cepellotti, Riccardo Sabatini, Nicola Marzari, and Boris Kozinsky. AiIDA: automated interactive infrastructure and database for computational science. Computational Materials Science, 111:218–230, 2016.
13Alessandro Genova. Davide Ceresoli, Alisa Krishtal, Oliviero Andreussi, Robert A. DiStasio Jr, and Michele Pavanello. eqe: An open-source density functional embedding theory code for the condensed phase. International Journal of Quantum Chemistry, 117(16):e25401, 2017.
14Jeongnim Kim, Andrew D Baczewski, Todd D Beaudet, Anouar Benali, M Chandler Bennett, Mark A Berrick, Nick S Blunt, Edgar Josué Landínez Borda, Michele Casula, David MCe pereley, Simone Chiesa, Bryan K Clark, Raymond C. Clay, Kris T Delaney, Mark Dewing, Kenneth P. Eler, Hongxia Hao, Oile Heineon, Paul R C Kent, Jaron T Krogel, Ilka Kylanpä, Ying Wu Li, M Grahame Lopez, Ye Luo, Fionn D Malone, Richard M Martin, Amrita Mathuraya, Jeremy McMinis, Cody A Melton, Lubos Mitas, Miguel A Morales, Eric Neuscamman, William D Parker, Sergio D Pineda Flores, Nichols A Romero, Brenda M Rubenstein, Jacqu-
line A R Shea, Hyeondeok Shin, Luke Shulenburg, Andreas F Tillack, Joshua P Townsend, Norm M Tubman, Brett Van Der Goetz, Jordan E Vincent, D ChangMo Yang, Yubo Yang, Shuai Zhang, and Luning Zhao. QMCPACK: an open source ab initio quantum Monte Carlo package for the electronic structure of atoms, molecules and solids. *Journal of Physics: Condensed Matter*, 30(19):195901, apr 2018.

D. Sangalli, A. Ferretti, H. Miranda, C. Attaccalite, I. Marri, E. Cancucia, P. Melo, M. Marsili, F. Paleari, A. Marrazzo, G. Prandini, P. Bonfà, M. O. Atambo, F. Affinito, M. Palumbo, A. Molina-Sánchez, C. Hogan, M. Grünning, D. Varzano, and A. Marinì. Many-body perturbation theory calculations using the yambo code. *Journal of Physics: Condensed Matter*, 31(32):325902, may 2019.

Jack Deslippe, George Samsonidze, David A. Strubbe, Manish Jain, Marvin L. Cohen, and Steven G. Louie. BerkeleyGW: A massively parallel computer package for the calculation of the quasiparticle and optical properties of materials and nanostructures. *Computer Physics Communications*, 183(6):1269 – 1289, 2012.

M. Govoni and G. Galli. Large Scale GW Calculations. *J. Chem. Theory Comput.*, 11:2680–2696, January 2015.

M. Schlüpf, H. Lambert, N. Zibouche, and F. Giustino. SternheimerGW: a program for calculating GW quasiparticle band structures and spectral functions without unoccupied states. *Computer Physics Communications*, 247:106856, 2020.

Giovanni Pizzini, Valerio Vitale, Ryotaro Arita, Stefan Blügel, Frank Freimuth, Guillaume Géranton, Marco Giberti, Dominik Gresh, Charles Johnson, Takashi Koresuene, Julen Ibañez-Azpíroz, Hyungjun Lee, Jae-Mo Lihn, Daniel Marchand, Antimo Marrazzo, Yuriy Mokrousov, Jakob M. Mustafa, Yoshiro Nohara, Yusuke Nomura, Lorenzo Paulatto, Samuel Poncè, Thomas Ponweiser, Junfeng Qiao, Florian Thöle, Stepan S Tisirkın, Małgorzata Wierzbowska, Nicola Marzari, David Vanderbilt, Ivo Souza, Arash A Mostofi, and Jonathan R Yatess. Wannier90 as a community code: new features and applications. *Journal of Physics: Condensed Matter*, 32(16):165902, jan 2020.

S. Poncè, E.R. Margine, C. Verdi, and F. Giustino. Epw: Electron-phonon coupling, transport and superconducting properties using maximally localized wannier functions. *Computer Physics Communications*, 209:116 – 133, 2016.

A. Otero de-la Roza, Erin R. Johnson, and Victor Luñáa. Critic2: A program for real-space analysis of quantum chemical interactions in solids. *Computer Physics Communications*, 185(3):1007 – 1018, 2014.

A. Ferretti, A. Calzolari, B. Bonferroni, and R. Di Felice. Maximally localized wannier functions from PAW or ultrasoft pseudopotentials. *J. Phys.: Condens. Matter*, 19:036215, 2007.

L. Calderín, V.V. Karasiev, and S.B. Trickey. Kubo-Greenwood electrical conductivity formulation and implementation for projector augmented wave datasets. *Computer Physics Communications*, 221:118 – 142, 2017.

Top500 supercomputers, 2019. https://www.top500.org/project/.

Michael Feldman. New GPU-Accelerated supercomputers change the balance of power on the TOP500, June 2018. https://www.top500.org/news/new-gpu-accelerated-supercomputers-change-the-balance-of-power-on-the-top500.

MAX: Materials at the eXascale. An EU Centre of Excellence for Supercomputing Applications. https://www.max-centre.eu.

H. Carter Edwards, Christian R. Trott, and Daniel Sunderland. Kokkos: Enabling manycore performance portability through polymorphic memory access patterns. *Journal of Parallel and Distributed Computing*, 74(12):2502 – 3216, 2014. Domain-Specific Languages and High-Level Frameworks for High-Performance Computing.

Erik Zenker, Benjamin Worpitz, René Widera, Axel Huelb, Guido Juckeland, Andreas Knüpfer, Wolfgang E. Nagel, and Michael Bussmann. Alpak-a - An Abstraction Library for Parallel Kernel Acceleration. IEEE Computer Society, May 2016.

A. Matthes, R. Widera, E. Zenker, B. Worpitz, A. Huelb, and M. Bussmann. Tuning and optimization for a variety of many-core architectures without changing a single line of implementation code using the alpak library. Jun 2017.

Rich Hornung, Holger Jones, Jeff Keasler, Rob Neely, Olga Pearce, Si Hammood, Christian Trott, Paul Lin, Courtenay Vaughan, Jeanine Cook, Rob Hoekstra, Ben Bergen, Josh Payne, and Geoff Womeldorff. ASC Tri-lab Co-design Level 2 Milestone Report 2015, Report No. LLNL-TR-677453, 2015.

ESL – the electronic structure library. https://esl.cccomp.org.

Electronic Structure Library coding Workshop: Drivers. Trieste, July 10-21 2017. https://gitlab.e-cam2020.eu/esl/ESLW_Drivers.

S. de Gironcoli. Lattice dynamics of metals from density-functional perturbation theory. *Phys. Rev. B*, 51:6773(R), 1995.

A. Dal Corso, A. Pasquarello, and A. Baldereschi. Density-functional perturbation theory for lattice dynamics with ultrasoft pseudopotentials. *Phys. Rev. B*, 56:R11369(R), 1997.

A. Dal Corso and S. de Gironcoli. Ab initio phonon dispersions of Fe and Ni. *Phys. Rev. B*, 62:273, 2000.

A. Dal Corso. Density-functional perturbation theory with ultrasoft pseudopotentials. *Phys. Rev. B*, 64:235118, 2001.

A. Dal Corso. Density functional perturbation theory for lattice dynamics with fully relativistic ultrasoft pseudopotentials: Application to fcc-Fe and fcc-Au. *Phys. Rev. B*, 76:054308, 2007.

A. Dal Corso. Density functional perturbation theory within the projector augmented wave method. *Phys. Rev. B*, 81:075123, 2010.

A. Floris, S. de Gironcoli, E. K. U. Gross, and M. Cococcioni. Vibrational properties of MnO and NiO from DFT+U-based Density Functional Perturbation Theory. *Phys. Rev. B*, 84:161102(R), 2011.

R. Sabatini, E. Küçükbenli, C.H. Pham, and S. de Gironcoli. Phonons in nonlocal van der Waals density functional theory. *Phys. Rev. B*, 93:235210, 2016.

E. T. Scholier, M. Calandra, and F. Mauri. Density functional perturbation theory for gated two-dimensional heterostructures: Theoretical developments and application to flexural phonons in graphene. *Phys. Rev. B*, 96:075448, 2017.

A. Urru and A. Dal Corso. Density functional perturbation theory for lattice dynamics with fully relativistic ultrasoft pseudopotentials: The magnetic case. *Phys. Rev. B*, 100:045115, 2019.

A. Floris, I. Timrov, B. Himmetoglu, N. Marzari, S. de Gironcoli, and M. Cococcioni. Hubbard-corrected density functional perturbation theory with ultrasoft pseudopotentials. *Phys. Rev. B*, 101:064305, 2020.

S. Baroni and R. Gebauer. The Liouville-Lanzcos Approach to Time-Dependent Density-Functional (Perturbation) Theory. Ref. 91, chapter 19, p. 375-390.

B Walker, A M Saïta, R Gebauer, and S Baroni. Efficient approach to time-dependent density-functional perturbation theory for optical spectroscopy. *Phys. Rev. Lett.*, 90(11):13001, 2006.

D Rocca, R Gebauer, Y Saad, and S Baroni. Turbo charging time-dependent density-functional theory with Lanzcos chains. *J. Chem. Phys.*, 128(15):154105, 2008.

Osman Barış Malcioglu, Ralph Gebauer, Dario Rocca, and Stefano Baroni. turbodDFT – A code for the simulation of molecular spectra using the Liouville–Lanzcos approach to time-dependent density-functional perturbation theory. *Comput. Phys. Commun.*, 182(8):1744–1754, aug 2011.

X. Ge, S.J. Binnie, D. Rocca, R. Gebauer, and S. Baroni. turbodDFT 2.0 — Hybrid functionals and new algorithms within time-dependent density-functional perturbation theory. *Comput. Phys. Commun.*, 185:2080–2089, 2014.

I. Timrov, O. Andreussi, A. Biancardi, N. Marzari, and S. Baroni. Self-consistent continuum solvation for optical absorption of complex molecular systems in solution. *J. Chem. Phys.*, 142:034111, 2015.

I. Tirumov, Nathalie Vast, Ralph Gebauer, and Stefano Baroni. Electron energy loss and inelastic x-ray scattering across sections from time-dependent density-functional perturbation theory. *Phys. Rev. B*, 88(6):064301, aug 2013. ibid. 91, 139901(E) (2015).

I. Tirumov, Nathalie Vast, Ralph Gebauer, and Stefano Baroni. turboEELS—A code for the simulation of the electron energy loss and inelastic X-ray scattering spectra using the Liouville–Lanzcos approach to time-dependent density-functional perturbation theory. *Comput. Phys. Commun.*, 196:460 – 469, 2015.

O. Motorny, N. Vast, I. Tirumov, O. Basseggio, S. Baroni, and A. Dal Corso. Electron energy loss spectroscopy of bulk gold with ultrasoft pseudopotentials and theLiouville-Lanzcos method. *Phys. Rev. B*, 102:035156, 2020.

Tommaso Gorni, I. Tirumov, and Stefano Baroni. Spin dynamics from time-dependent density-functional perturbation theory. *The European Physical Journal B*, 91(10), 2018.

https://www.w3.org/standards/xml/schema for XML schema defi-
nition (XSD).

https://pypi.org/project/xmlschema/ for the xmlschema package.

https://gitlab.com/QEF/xmltool for the xmltool package.

https://pypi.org/project/geschema/ for the geschema package.

Ask Hjorth Larsen, Jens Jørgen Mortensen, Jakob Blomqvist, Ivan E. Castelli, Rune Christensen, Marcin Dulak, Jesper Friis, Michael N. Groves, Bjørst Hammer, Cory Hargus, Eric D. Hermes, Paul C. Jennings, Petter Bjørre Jensen, John K. Ketchin, Esben Leonhord Kolsbjerg, Joseph Kubal, Kristen Kaasbjerg, Steen Lyssgaard, Jön Bergmann Maronsson, Tristan Maxson, Thomas Olsen, Lars Pastewka, Andrew Peter- sen, Carsten Rostgaard, Jakob Schiøtz, Ole Schütz, Mikkel Strange, Kristian S. Thygesen, Tjejs Vege, Lasse Vilhelmsen, Michael Walter, Zhehuna Zeng, and Karsten Wedel Jacobsen. The atomic simulation environment—a python library for working with atoms. *J. Phys.: Condens. Matter*, 29, 2017.

Shyue Ping Ong, William Davidson Richards, Anubhav Jain, Geoffroy Hautier, Michael Kocher, Shreyas Cholia, Dan Gunter, Vincent L. Chevrier, Kristin A. Persson, and Gerbrand Ceder. Python materials genomics (pymatgen): A robust, open-source python library for materials analysis. *Computational Materials Science*, 68:314 – 319, 2013.

Anton Kokalj. XyCeSdn—a new program for displaying crystalline structures and electron densities. *Journal of Molecular Graphics and Modelling*, 17(3):176 – 179, 1999.

https://github.com/QEF/postqe for the postqe package.

Filippo Spiga and Ivan Girotto. phGEMM: A CPU-GPU Library for Porting QUANTUM ESPRESSO on Hybrid Systems. In 2012 20th Euromicro International Conference on Parallel, Distributed and Network-based Processing. IEEE, February 2012.

Filippo Spiga. QE-GPU: GPU-Accelerated Quantum ESPRESSO, 2012-2017.

Ivan Girotto, Nicola Varini, Filippo Spiga, Carlo Cavazzoni, Davide Ceresoli, Layla Martin-Samos, and Tommaso Gorni. Enabling of QUANTUM ESPRESSO to Petascale Scientific Challenges. PRACE, 2012.

Joshua Romero, Everett Phillips, Gregory Ruetsch, Massimiliano Fatica, Joshua Romero, Everett Phillips, Gregory Ruetsch, Massimiliano Fatica, Filippo Spiga, and Paolo Giannozzi. A performance study of QUANTUM ESPRESSO’s PWscf code on multi-core and GPU systems. In Internal- tional Workshop on Performance Modeling, Benchmarking and Simulation of High Performance Computer Systems, pages 67–87. Springer, 2018.

GPU-accelerated QUANTUM ESPRESSO 11 Zenodo. http://doi.org/10.5281/zenodo.823200.

Optionally, cuRAND can also be used for the generation of random wave- functions.

Carlo Cavazzoni, Fabio Affinito, Uliana Alekseeva, Claudia Cardoso, Augustin Degomme, Pietro Delugas, Andrea Ferretti, Alberto Garcia, Luigi Genovese, Paolo Giannozzi, Anton Kozhevnikov, Ivan Marri, Stephen Mohr, and Daniel Wortmann. First report on code profiling and bottleneck identification, structured plan of forward activities. Deliverable D4.2 of the H2020 CoE MAX. EC grant agreement no: 824143, CINECA, Bologna, Italy (2019). Available at max-centre.eu/project-repository.

Daniel Wortmann, Uliana Alekseeva, Stefano Baroni, Augustin Degomme, Pietro Delugas, Stefano de Gironcoli, Andrea Ferretti, Alberto Garcia, Luigi Genovese, Paolo Giannozzi, Anton Kozhevnikov, and Ivan Marri. First release of MAX software: report on the performance portability. Deliverable D2.1 of the H2020 CoE MAX. EC grant agreement no: 824143, Julich, Germany (2019). Available at max-centre.eu/project-repository.

QUANTUM ESPRESSO input generator and structure visualizer, https://www.materialscloud.org/work/tools/qeinputgenerator.

Gianluca Prandini, Antonio Marrizzo, Ivan E. Castelli, Nicolas Mounet, and Nicola Marzari. Precision and efficiency in solid-state pseudopotential calculations. *npj Computational Materials*, 4(1), December 2018.

Alex Willand, Yaroslav O. Kvasnhin, Luigi Genovese, Álvaro Vázquez-Mayagoitia, Arpan Krishna Deb, Ali Sadeghi, Thierry Deutsch, and Stefan Goedecker. Norm-conserving pseudopotentials with chemical accuracy compared to all-electron calculations. *The Journal of Chemical Physics*, 138(10):104109, March 2013.

Martin Schlüpf and François Gygi. Optimization algorithm for the generation of ONCV pseudopotentials. *Computer Physics Communications*, 196:36–44, November 2015.

M.J. van Setten, M. Giantomassi, E. Bouquet, M.J. Verstraete, D.R. Hamann, X. Gonze, and G.-M. Rignanese. The PseudoDojo: Training and grading a 85 element optimized norm-conserving pseudopotential datafile. *Computer Physics Communications*, 226:39–54, May 2018.

Kevin F. Garrity, Joseph W. Bennett, Karin M. Rabe, and David Vanderbilt. Pseudopotentials for high-throughput DFT calculations. *Computational Materials Science*, 81:446–452, January 2014.

Andrea Dal Corso. Pseudopotentials periodic table: From H to Pu. *Computational Materials Science*, 95:337–350, December 2014.

M. Topsakal and R.M. Wentzovich. Accurate projected augmented wave (PAW) datasets for rare-earth elements (RE=E-La–Lu). *Computational Materials Science*, 95:263–270, December 2014.

Miguel Quiróis, Saulius Gražulis, Saule Girdzijauskaite, Andrius Merkys, and Antanas Vaitkus. Using SMILES strings for the description of chemical connectivity in the Crystallography Open Database. *Journal of Chemical Informatics*, 10(1), May 2018.

Andrius Merkys, Antanas Vaitkus, Justas Butkus, Mykolas Okuličiūnas, Visvaldas Kairys, and Saulius Gražulis. COD::CIF::Parser: an error-correcting CIF parser for the Perl language. *Journal of Applied Crystallography*, 49(1), Feb 2016.

Saulius Gražulis, Andrius Merkys, Antanas Vaitkus, and Mykolas Okuličiūnas. Computing stoichiometric molecular composition from crystal structures. *Journal of Applied Crystallography*, 48(1):85–91, Feb 2015.

Saulius Gražulis, Adriana Daškevič, Andrius Merkys, Daniel Chatteiginer, Luca Lutterotti, Miguel Quirós, Nadezhda R. Serebrayana, Peter Moeck, Robert T. Downs, and Armel Le Bail. Crystallography Open Database (COD): an open-access collection of crystal structures and platform for world-wide collaboration. *Nucleic Acids Research*, 40(D1):D420–D427, 2012.

Saulius Gražulis, Daniel Chatteiginer, Robert T. Downs, A. F. T. Yokochi, Miguel Quirós, Luca Lutterotti, Elena Manakova, Justas Butkus, Peter Moeck, and Armel Le Bail. Crystallography Open Database – an open-access collection of crystal structures. *Journal of Applied Crystallography*, 42(4):726–729, Aug 2009.

R. T. Downs and M. Hall-Wallace. The American Mineralogist Crystal Structure Database. *American Mineralogist*, 88:247–250, 2003.

Paolo Giannozzi, Oscar Baseggio, Pietro Bonfà, Roberto Car, A. Rubio, editors. *Pseudopotentials for high-throughput DFT calculations*. *Computer Physics Communications*, 198:225–287, 2015.

Kevin F. Garrity, Joseph W. Bennett, Karin M. Rabe, and David Vanderbilt. Pseudopotentials for high-throughput DFT calculations. *Computational Materials Science*, 81:446–452, January 2014.