Zeitschrift: Helvetica Physica Acta

Band: 61 (1988)

Heft: 1-2

Artikel: Common elements of the various disciplines in computational science

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DOI: https://doi.org/10.5169/seals-115932

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COMMON ELEMENTS OF THE VARIOUS DISCIPLINES

IN COMPUTATIONAL SCIENCE

Talk at Symposium on Computational Physics Swiss Physical Society, Fall Meeting 1987

R. Henzi, University of Berne

ABSTRACT

The advent of parallel computers will have an important influence on many areas of science, just as have other new tools, such as particle accelerators, neutron sources, large telescopes, or space labs. Its great advantage is its general applicability to all branches of science. The computational scientist is faced with fundamentally new challenges because the high rate of computation is only fully realized if the algorithms and programming techniques are tailored to the architecture of the computer. This is illustrated by an example of vectorization and optimization of an IF-loop. Two classes of examples are described where important new scientific or engineering results are being obtained by large-scale computer simulation: simulation of classical or pseudo-classical many-body systems, and of statistical and quantum systems. Common computational techniques in these classes are described. Many other examples of application areas of supercomputers are given, and the case is made that in the computational approaches to these pressing and challenging problems the mathematical and computational aspects cannot and should not be considered in isolation. A new forum of collaboration of scientists in Switzerland, SPEEDUP, It forms an interface between is presented. advanced computers, such as parallel computers, and the physicist or engineer with the problem.

1. INTRODUCTION

Current nominal computational performance rates of the types of supercomputers used in large-scale simulations in science and engineering are of the order of 100s of MFLOPS (an MFLOP is 1 million floating point operations per second). The supercomputers available during the early 1990s will have rated performances in the 10,000 MFLOP range. The mid 1990s machines are forecast to be 300,000 MFLOP machines, and those of the turn of the millenium should be over 1,000,000 MFLOPs in performance. (Figure 1.)

These computers obtain their speed principally by an increased use of what is called PARALLELISM, more precisely, vector and array parallelism, and are collectively termed parallel computers.

Applications on machines of such enormous power will be themselves tremendously large software systems developed over many years by organized groups of scientists and engineers.

The advent of such powerful computational tools will have an important influence on many areas of science, just as have other new tools, such as particle accelerators, neutron sources, large telescopes, or space labs. (Section 2.) Its great advantage is its general applicability to all branches of science.

In comparison with his work on the classical serial "FORTRAN machines", the computational scientist is faced with fundamentally new challenges because the high rate of computation is only fully realised if the algorithms and programming techniques are tailored to the architecture of the computer. (Section 3.)

To show the types of scientific problem to which parallel computers can profitably be applied, we describe two classes of The first class is the simulation of classical or pseudo-classical many-body systems, where we find examples such as the simulation of spiral structure in galaxies of stars, the simulation of the flow of electrons in semiconducting devices, and the simulation of melting and glass formation in ionic materials. The second class is the simulation of statistical and quantum systems, where we find examples such as the simulation of a highly idealized model, e.g. the Ising me equilibrium or far from equilibrium, or the simulation of the Ising model in lattice gauge theories, the simulation of continuum and lattice models of polymers, and the simulation of atomic models of quantum liquids and solids. For each of these two classes a common computational technique (the particle-mesh method in the first class, the Monte Carlo method in the second class) has been developed and is used in all the diverse applications. (Sections 4 and 5). This makes apparent the importance of supporting research in Computational Science, as an activity in its own right, in order to develop methods suited to the new parallel computers.

We have only given two classes of examples where a parallel computer can have an important effect on the advancement of science and engineering. Other examples abound in e.g. hydrodynamic systems, chemical systems and combustion, plasma



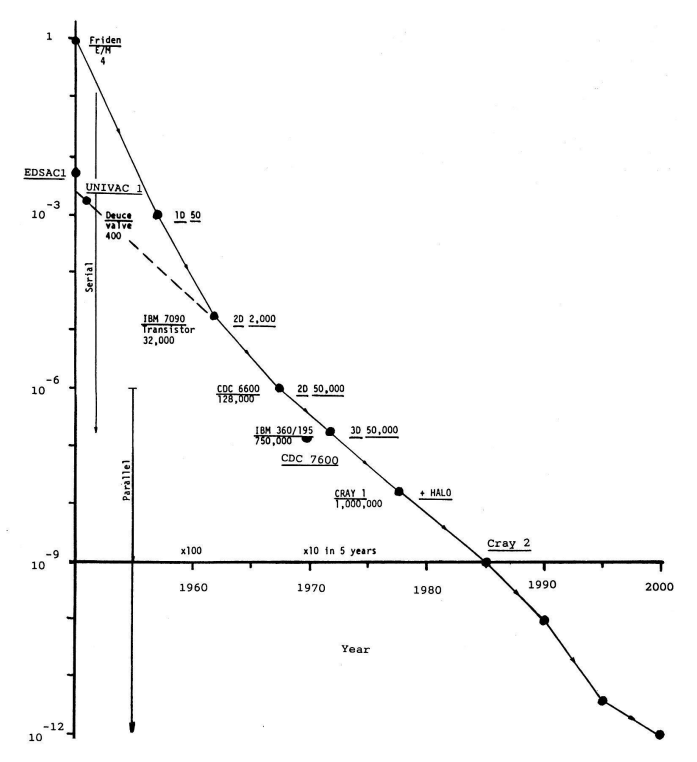


Figure 1. History (since 1950) and Forecast of Computer Arithmetic Speed.

physics, geophysical applications, meteorology, structural mechanics, nondestructive testing and tomographic reconstruction, and mathematical models in the biological sciences. In the computational approaches to these pressing and challenging problems the mathematical and computational aspects cannot and should not be considered in isolation. (Section 6).

A Cray 1 was installed at the Swiss Federal Institute of Technology Lausanne in early 1986. A national initiative for supercomputing in Switzerland [Hochstrasser 88] was voted by Swiss Parliament in Summer of 1986. Early installation of more powerful supercomputers is also necessary in this framework and is eagerly awaited. Currently, a transitory solution is being considered in which a Cray X-MP at the Swiss Federal Institute of Technology Zurich and a Cray 2 at the Swiss Federal Institute Lausanne would bridge the time till a supercomputer of the latest generation becomes available.

Vast amounts of computer power will be available for users of the new Swiss supercomputer system. Effective distribution of these resources presupposes the use of

- advanced, high level, high-bandwidth communication facilities, and
- a very large user base.

Support of this large user base also presupposes the maintenance of a substantial infrastructure of general purpose computer facilities providing

- job preparation,
- administration, and
- general computer service support.

It is likely that such an infrastructure would have to be distributed in structure, if only because of the complexity of the service environment.

The real question we face is: how should this entire supercomputing environment be conceived to turn raw computer power into advances in human knowledge and engineering? (Section 7.)

2. THE INFLUENCE OF COMPUTER SPEED ON THE COMPLEXITY OF PHYSICAL PROBLEMS THAT CAN BE SOLVED

Figure 1 shows the time to perform a floating-point multiply on a number of computers since 1950, together with the size of high-speed memory, and the above-metioned forecast to the year In recent years there has been an increase of a factor of 10 in speed every 5 years. This has been obtained partly by increases in the speed of logic elements and partly by the introduction of more parallelism in the architecture of the computer.

Let us consider that a computer run of a few hours is the maximum a single user can reasonably perform on an installation. Then, in the study of galactic evolution, the first valve computers would have been limited to the study of about 50 stars in a rather unrealistic one-dimensional galaxy. The first transistor computers, such as the IBM 7090, allowed the first realistic simulations of 2000 stars moving in a plane. The CDC 6600 allowed the number of stars to be increased to 50 000. But the move to full three-dimensional motion had to await the arrival of the CDC 7600 and IBM 360/195. In the above simulations only the disk stars are simulated and the halo stars, that lie out of the plane of the galactic disc, are represented as a fixed external gravitational field. The opportunity to represent moving stars in the halo requires the extra computer speed of the parallel computer, as exemplified by the Cray 1 computer.

3. PROGRAMMING VECTOR AND OTHER PARALLEL COMPUTERS AND ITS IMPLICATION FOR INTERDISCIPLINARY GROUPINGS

In large-scale simulation codes, the calculation is best performed entirely in high-speed memory. The calculation time is determined primarily by the time to perform floating-point arithmetic because in any carefully written code accesses to and from memory, index calculations and house-keeping calculations can be performed in parallel with the floating-point arithmetic. Provision for doing this exists on most top-of-the-range machines (e.g. IBM 3090, Crays, Cybers, Fujitsus). Accordingly, such organizational operations may be ignored and the performance of computers may be measured in terms of the number of millions of floating-point operations performed per second. For this purpose, all floating-point additions, subtractions, multiplications and divisions are counted with equal weight. (To distinguish separately between the different times for these operations would complicate matters without changing the conclusions much.)

Normally, any standard ANSI FORTRAN 77 program can be executed on a vector computer, such as the Cray 1, Cray X-MP, Cray 2, Cyber 205, ETA 10 or Fujitsu, through a compiler which recognizes DO-loops that can be executed by the explicit vector instructions of the machine. Existing FORTRAN programs, which were written without knowledge of the machine architecture, will in many cases run faster than on the IBM 3090. But remember, running a program on a vector processor does not necessarily mean that one uses the vector instructions of the machine - the real power of these new computers. The full vector speeds, in terms floating point operations per second, of 5 to 25 times that of the scalar speed can be obtained in FORTRAN provided simple rules, designed to permit the compiler to vectorize, are obeyed. This increased performance gain results in increased motivation to perform the optimization of programs for these machines by applying these rules.

Another issue which should motivate the programmer is that vector and some multi-processors are here to stay in scientific computing, and we will probably even see them in other areas of computing, e.g. transaction processing. The vector processor is not a passing fad, and the scientific computer programmer should learn how to use it.

As examples of the type of vectorization and optimization work

referred to above, we mention three that have recently been completed within the framework of the PARCOM Project on Scientific Applications and Algorithm Design for Parallel Computing at the University of Berne. These involve Monte Carlo updating [Decker and Henzi 87; Decker 87], problems to test parallel computers [Henzi 88a], and inverse problems [Henzi 88b].

For illustration, let us vectorize an example which used to be considered non-vectorizable. All too often people cop out on a little work by saying, "This code isn't conductive to vectorization." In this IF-loop there is a good calculation we would like to vectorize:

```
M=0
100 CONTINUE
         M=M+1
         A(M)=B(M)**2 + .5*C(M)*D(M)/E(M)
     IF (A(M).GT.O) GO TO 100
```

The problem we have is that we are not sure when the IF-loop, whose timing is illustrated in Fig. 2a, is terminated. Split the calculation out from the decision:

```
DO 21 M=1,N
         TEMP(M)=B(M)**2 + .5*C(M)*D(M)/E(M)
21
     CONTINUE
     M=0
100
     CONTINUE
         M=M+1
         A(M) = TEMP(M)
     IF (A(M).GT.O) GO TO 100
```

This restructuring will not run faster on a scalar machine; however, it should on a vector computer. Of course, as is illustrated in Fig. 2b, a lot depends on which value of A is the first not to satisfy the test. Quite often a DO-loop may vectorize; however, it could be written to vectorize even better. On the Cray we get best performance for a vector length Therefore, we stripmine this example and use the ILLZ function from SCILIB to find the first occurrence where the A array is less than zero:

```
DO 21
            I=1, N, 64
         LENGTH=MINO(64, N-I+1)
         DO 20 MM=1, LENGTH
             M=I+MM-1
              TEMP(MM) = (B(M)**2+.5*C(M)*D(M)/E(M))-1.E-1000
20
         CONTINUE
         NTOP=ILLZ(LENGTH, TEMP, 1)
         DO 31 MM=1,NTOP
             M=I+MM-1
              A(M) = TEMP(MM)
31
         CONTINUE
         IF(NTOP.LT.LENGTH) GO TO 22
21
     CONTINUE
22
     CONTINUE
```

The corresponding timing is illustrated in Fig. 2c. Figures 2a-c also illustrate that the performance gain due to a code

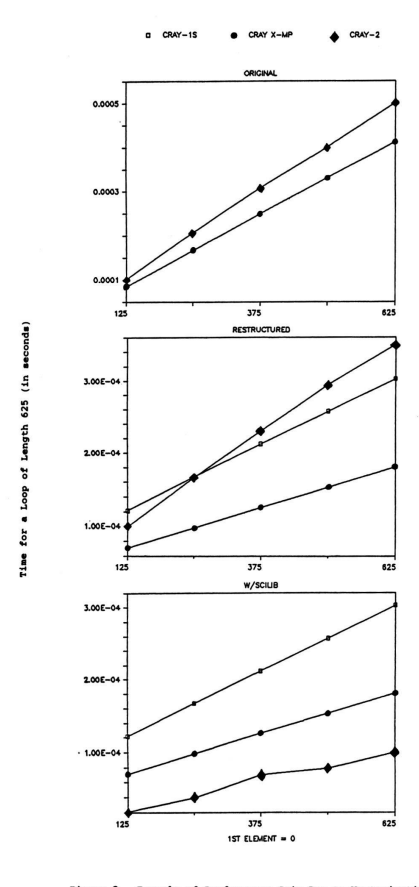


Figure 2. Example of Performance Gain Due to Vectorization and Optimization.

(a) Original, (b) Restructured, (c) With SCILIB.

restructuring can not only be quite machine dependent but also quite considerable, as e.g. for the Cray 2. On the one hand, this makes apparent that optimization can be profitable. On the other hand, optimization of a code for large-scale simulation can entail a considerable investment of human resources. But then it becomes interesting to exploit such an optimized code by applying it to simulate as many different physical systems as possible, optimizing in this way that investment. This in turn favors the formation of a computational community in the form of an interdisciplinary grouping based on the common underlying mathematics and algorithmics of research areas. Similar considerations also apply to computers incorporating array parallelism, either without or in addition to vector parallelism.

4. SIMULATION OF CLASSICAL OR PSEUDO-CLASSICAL MANY-BODY SYSTEMS

Three principal kinds of simulation of such systems may be identified: particle-particle simulation, e.g. molecular dynamics simulation to study the structure of liquids, or simulation of stellar clusters, or of clustering of galaxies; particle-mesh simulation, e.g. galaxy simulation, or simulation of dilute plasmas, or of microscopic semiconductor devices; and particle-mesh/particle-particle simulation, e.g. simulation of phase changes in ionic solids and liquids (such as KC1, CaO), or of clustering of galaxies, or of a two-dimensional electron film.

For the first kind, the action-at-a-distance formulation of the force law is used. For the second kind, the force is treated as a field quantity approximating it on a mesh. The third kind is a hybrid of the former two. Partly the physics of the phenomenon under investigation and partly the consideration of the computational cost determine the choice. Pioneering work, especially on particle-mesh simulation, has been done by Hockney and Eastwood [1981] and their collaborators.

To illustrate these kinds of simulation, let us now consider a particle-mesh/particle-particle simulation of an ionic solid/liquid system. The long-range interaction is of Coulomb type. It may be found by accumulating the charges into a charge density as a source and computing the electrostatic potential from Poisson's equation. The long-range force on any particle is then found by computing the gradient of the potential at the particle's position. In addition, pair-wise short-range forces from nearby particles have to be included to take into account the repulsion of the electronic shells.

The system is completely described by storing the positions and velocities of all the particles in the computer. To solve Poisson's equation, a computational mesh is introduced. During the charge assignment phase, first, a given number of neighbouring mesh points of a particle are found. Then the charge is assigned to them so that computational errors are minimized for the mesh points used, or equivalently to obtain the best quality/cost compromise. Different assignment schemes arise by the choice of different numbers of neighbouring points.

The state of the physical system at some time t is described by

the set of particle position and velocity vectors xi(t), vi(t), where $i=1,\ldots,N$. The timestep loop updates these values using the forces of interaction and equations of motion to obtain the state of the system at a slightly later time t+DT, as follows:

C Build up charge distribution rho(x)

LOOP FOR all particles

Find neighbouring mesh points of particle

Assign charge to mesh

ENDLOOP

C Solve Poisson's equation on mesh to obtain potential phi(x)

LOOP FOR all mesh points

Solve divgrad*phi = - rho/epsilon0

ENDLOOP

C Compute forces and accelerate each particle of mass m for C time step DT by changing particle velocity according to

C total force

LOOP FOR all particles

Find mesh force FM on particle

Find short-range force FS from near-neighbour particles

vi(new) := vi(old) + (FM+FS)*DT/m

ENDLOOP

C Move each particle to new position

LOOP FOR all particles

xi(new) := xi(old) + vi(new)*DT

ENDLOOP

The position and velocities of all particles are now known at the new time-level. Repeated application of the time-step loop allows to follow the evolution of the system in time. For a system without short-range forces FS is omitted. The point of this algorithm is that it has an arithmetic operation count proportional to N, and not to N*N as would be the case if all N*N pair interactions were evaluated directly. Furthermore, Fast Fourier Transform techniques are used to solve Poisson's equation.

A galaxy simulation is done by changing the sign in Poisson's equation, and replacing the specific charge of a particle by the

square root of the gravitational constant G. Submicron semiconductor simulation differs from this scheme in that important short time-scale effects due to electron scattering by the crystal lattice must be taken into account. This is done by replacing the mobility approximation, valid in devices whose dimensions are large compared with the electron mean free path, by a microscopic description in which statistical Monte Carlo methods are used to simulate the scattering of the electrons by the crystal lattice.

5. SIMULATION OF STATISTICAL UND QUANTUM SYSTEMS

In the simulation of statistical systems in physics, the equation of state, transport coefficients (pressure, viscosity, and thermal conductivity, for example), and other macroscopic properties of matter are related to and derived from intermolecular forces. The area has diverse and important applications, ranging from metallurgy to polymer chemistry to semiconductors. It is an active area of research from the points of view of theory, numerical methods, and experiment. One numerical method for statistical physics is basically of the type of molecular-dynamics calculations already mentioned in the previous section.

The equations of statistical physics involve a large or infinite number of degrees of freedom. Hence, the mathematical theory of use here is the analysis over infinite-dimensional spaces.

In the study of quantum fields, almost the same mathematical structure arises, the relationship between the two being that a quantum field is a continuum limit of a statistical physics (crystal or lattice) model [Wilson 74]. This analogy is illustrated by the equations for the equilibrium distribution $d\omega$ and the partition function Z:

$$d\omega = e \prod_{i} dx_{i}, i=1,...,N$$

$$d\omega = e \prod_{i} d\phi_{x}$$

$$Z = \int d\omega$$

$$Z = \int d\omega$$

$$V = \sum_{i < j} U(x_{i}-x_{j})$$

$$S = \int L(\phi_{x}, \partial \phi_{x}/\partial x) d^{4}x$$

where V is the intermolecular potential energy taken as a sum of pair potentials U, and S is the classical action taken as a space-time integral of the Lagrange density depending on the

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quantum field $\Phi_{\mathbf{x}}$.

Important numerical methods are the Monte Carlo methods used in:

- the quadrature of very-large-dimensional spaces, such as the determination of $d\omega$ above, as well as in

 the direct simulation of stochastic systems, an example of which we have briefly discussed at the end of the previous section.

These methods have become an "experimental" tool of theoretical physics. In studying the qualitative and quantitative behaviour of the kind of systems described in the introduction, one can carry out numerical studies in which the high-dimensional (i.e., many-body) character of the problem is not distorted.

In the application to the calculation of the equilibrium distribution $d\omega$, the essence of the Monte Carlo method is as follows:

- Starting from an arbitrary point in a given ensemble, one modifies a single particle position x; "at random" but usually so as to lower the potential energy V, occasionally so as to raise V.
- After enough such elementary steps, convergence to the distribution $d\omega$ is obtained.

The procedure is analogous for quantum fields. As described here, the method is extremely simple, and complications arise from the necessity to obtain convergence in a reasonable time for realistic problems.

In statistical physics, perhaps the most significant success has been in the microscopic theory of fluids, where Monte Carlo simulation has provided the "experimental" basis for accurate series expansions in $\exp(-\beta U)-1$ of the expression for $d\omega$, which are especially useful to give weak corrections, such as real-gas corrections to an ideal gas. Other conspicuous successes are the extraction of critical exponents by joining the ideas of the renormalization group to Monte Carlo simulation. Monte Carlo simulation of a kinetic Ising model gives a quantitative account of scattering experiments on a real alloy, and Monte Carlo methods applied to the many-body nonrelativistic Schrödinger equation give a quantitative account of the energy of real liquid helium as a function of the density.

In quantum field theory the most significant successes have perhaps been in the determination of the quark-antiquark potential and of the nature of phase transitions, in studies on the existence of a renormalizable continuum theory and on the correct interpretation of lattice results, and in the computation of the "glue ball" spectrum. Examples of this type of work, in collaboration with the Institute for Theoretical Physics of the University of Berne, have recently been completed within the framework of the PARCOM Project already mentioned in section 3 [Carrupt et al. 1987; Decker et al. 1987].

6. UNITY AMONG THE VARIOUS STEPS OF THE OVERALL SIMULATION PROCESS

In the computational approaches to most of today's pressing and challenging scientific and technological problems the mathematical and computational aspects cannot and should not be considered in isolation. There is a unity among the various steps of the overall simulation process:

- from the formulation of the physical problem
- to the construction of appropriate mathematical models
- to the design of suitable numerical methods
- to their computational implementation, and, last but not least
- to the validation and interpretation of the computed results.

The steps are more often than not deeply interconnected and the computational process may indeed be part of the model construction. We may illustrate this by viewing the same problem from three different standpoints [Henzi 87a].

In Table 1 (Tabelle 1) the traditional approach is taken of looking at a number of scientific and engineeering disciplines leading to computational simulation from the viewpoint of the mathematical formulation of typical problems to which they give rise:

HSy Hydrodynamic Systems CSy Chemical Systems and Combustion PlPh Plasma Physics TPh Particles Physics PhM Condensed-Matter Physics Geophysical Applications GPh Met Meteorology Ast Astrophysical Applications StM Structural Mechanics 3D Nondestructive Testing and Tomographic Reconstruction (three-dimensional) Mathematical Models in the Biological Sciences BioM ElK Electronic Components

In Table 2 (Tabelle 2) these disciplines are touched upon again, this time from the viewpoint of the computational and mathematical difficulties that arise in connection with them. E.g., these difficulties may involve nonlinearities, large numbers of degrees of freedom, different scales of length and time, or singularities of various types.

In Table 3 (Tabelle 3) the viewpoint becomes that of the numerical algorithms involved in the computations, such as various discretization methods, continuation approaches, splitting techniques and Monte Carlo methods.

Of necessity, many topics have been left out. Some of these topics as well as some of those above have been considered more

	HSy	CSy	PIPh	TPh	PhM	GPh	Met	Ast	StM	3D	BioM	EIK
Partielle Differentialgleichungen												
Inverse Probleme, Integralgig. Minimalisierung unter Zwangsbed	ָם י											
Gewöhnliche Differentialgig.				u								0
Hamiltonsche Dynamik												
Gruppen und Quantenfeldtheorie Verteilungen, Zustandssummen												
Quadratur hochdimens. Räume		_		0				_				
Simulation stochastischer Syst. Informations-(Signal-)Verarb.												
Kombinatorik und Graphen												
Tabelle 1 Mathematische Form	ulien	ungen	ľ									
	HSy	CSy	PIPh	TPh	PhM	GPh	Met	Ast	StM	3D	BioM	EIK
Nichtlinearitäten								0				
Freiheitsgrade Versch. Zeit- & Längenskalen									0		0	0
Singularitäten in Koeff., Daten							_					
Randbedingungen Bifurkation, Chaos, SymmBrech	. 0				0						0	0
Schlechtgestellte inv. Probleme						0						
Eff. Medien, Homogenis., Randon Validierung, Fehlerabschätzung	ם							0			0	0
Tabelle 2 Numerische und math	nema	tische	Schw	ierig	keiten	i.						
	HSy	<u>CSy</u>	PIPh	TPh	PhM	GPh	Met	Ast	StM	30	BioM	EIK
Diskretisierungsmethoden:						Caracti				16.16		_
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Finite Differenzen Varitationsmethoden Finite Elemente Transformationsmethoden Semidiskrete Methoden Charakteristiken Adaptive Grid Methoden	0000	0	0	000	0	0	0	0	00	0	0	0
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Finite Differenzen Varitationsmethoden Finite Elemente Transformationsmethoden Semidiskrete Methoden Charakteristiken Adaptive Grid Methoden Meth. zur Lsg. diskretisierter Diff. Direkte Methoden Indirekte, Multigrid Methoden Fortsetzungs- und Homotop. M. Optimierungsmethoden Graphentheoretische Methoden Splitting M. & Defect Corr. Monte Carlo Techniken Problemabhängige Methoden: Riemann Probl. & Wellenmoder Asymptotic Analysis Front Tracking Wirbelmethode Skaleninv. & Renormierungsgr.	0000 00;	0 0 000 00 00	0 00 0	0000 0 000000	0 0 00 000	0	0	0 0 0 0 0 0 0 0	000 0 000	0	0	0

closely in an analysis of the importance of supercomputers for the telecommunications industry [Henzi 87b]. Neither was it attempted to address all computational and mathematical difficulties nor all variations of numerical algorithms.

7. TURNING RAW COMPUTER POWER INTO ADVANCES IN HUMAN KNOWLEDGE

From a computer modelling point of view, the unified framework provided e.g. by statistical mechanics, that was illustrated in Section 5, appears as the similarity of mathematical models and thence as the similarity of numerical algorithms and of computer programs for a wide range of physical phenomena. One benefit of the unified approach to a wide range of physical phenomena is that advances in computational methods in one area of application more rapidly find their way into others.

The principal benefit of considering computational science in general and e.g. simulation of many-body or of statistical and quantum systems in particular as entities appears when we look at the computer program. As we have seen, programs developed to study particular phenomena may, with minor modifications, be used to study something completely different. Large economies in human power and computer resources can be made by recognizing the similarities of physical, mathematical, numerical and computational problems, and by embodying the similarities in standardized structured programs. ASTRID, presented by Ralf Gruber [Gruber 88] at this symposium, is an example of such an approach.

Last year, the SPEEDUP organization was established in Switzerland with the objective of promoting vector and parallel computing in science and industry, as well as the interest that academic and industrial research workers always have at their disposal a system based on the current top-of-the-range number-crunching computer. The SPEEDUP activities have started out with the publication of the journal SPEEDUP, of which the first issue has appeared on November 1, 1987, and with the organization of workshops, of which the first took place on November 20, 1987, on the theme "Supercomputing - The State of the Art". Coming workshops in 1988 will take place as follows:

January 15	Parallel Computers
March 18	The ASTRID Project of EPF Lausanne: A Family of Simulation Programs Adapted to Parallel Vector Computers
May 20	Simulation of Quantum and Statistical Systems
September 16	Computational Quantum Chemistry (tentative)
November 18	The New National Supercomputing System for Switzerland (tentative)

Subscriptions to the SPEEDUP journal and requests for information should be mailed to: SPEEDUP Secretariat, IAM, Uni Bern, Länggassstrasse 51, CH-3012 Bern, Switzerland; Telephone 031/65 86 81; E-Mail u02e@cbebda3t.bitnet.

Henzi H.P.A.

Traditional subject divisions lead academia to teach experimentation and theory to the physicist, numerical analysis to mathematicians, and computer architecture to computer scientists, so that it is rare to find an individual who has the combination of backgrounds required for the type of work in large-scale simulation discussed here. SPEEDUP and, more generally, Computational Science form interfaces between advanced computers, such as parallel computers, and the physicist or engineer with the problem; these interfaces intend to help the research worker to find the combination of the best computer architecture, the best numerical algorithms and the computer simulation that can bring new insight and discoveries in science and make a contribution to the better design of industrial products.

ACKNOWLEDGEMENT

The support of the PARCOM Project by the Swiss National Science Foundation, the support of the author, through a Bilateral Exchange Grant, by the Swiss National Science Foundation and the Natural Sciences and Engineering Research Council Canada, and the hospitality of University of Berne and of ETH Zürich are gratefully acknowledged.

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