

'GUIDE: Groningen University interactive display of enzymes and other biological macromolecules' **Hol, W G J and Postma, J** Laboratory of Chemical Physics, University of Groningen, The Netherlands.

GUIDE is a program which runs on a PDP-11/34 which is the controller of an Evans and Sutherland Picture System 2. The PDP is connected by DECNET to a PDP-11/45, which in its turn is connected to a Cyber 170/760 and to a PDP-11/70. The PDP has access to one 205 Mbyte RA 80 disc. It is intensively used by the Groningen protein crystallography group for a variety of purposes. The capabilities of the program will be illustrated by examples taken from the research projects going on in the laboratory. Topics are:

- refinement of protein structures
- simultaneous investigation of several electron density maps plus a molecular model
- simultaneous display and manipulation of up to 4 different molecular models
- creating 'new' proteins by altering the amino acid side chains according to a new sequence

The striking similarity of dinucleotide-binding $\beta\alpha\beta$ -units in six different proteins will be described. The prediction of the 3D structure of a crucial part of an oncogenic protein will be highlighted.

'Application of a parallel processor to molecular graphics' **Hubbard, R, Fincham, D and Quinn, J E*** University of York. *Queen Mary College.

A disadvantage of many interactive molecular modelling packages is their inability to produce interactive space filling representations of the CPK type. These representations would allow a fuller appreciation of the spatial relationships present than the 'dot surface' types used on vector machines (making, for example, the interaction of enzymatic active sites with model substrates much clearer). In this paper an algorithm for drawing such models using a truly parallel processor (the ICL DAP (digital array processor) at Queen Mary College in London) is discussed along with its implementation using DAP Fortran to produce an image for display on a raster graphics device.

In general terms the algorithm is an application of the 'Z buffer' method but makes use of the very powerful masking techniques available on the DAP to calculate blocks of 64×64 pixels simultaneously. Although no formal optimization of the program has been carried out, use has been made of the variable word lengths (standard features on the DAP) wherever arithmetic is required, to provide substantial decreases in execution time. (Already we can create these pictures in times of about half a millisecond/atom).

A combination of the power of the parallelism of the DAP allied with any suitable graphical output device, will enable interactive computation of space filling models, including whole and part rotations of the molecule, thereby providing a significant increase in

the ability of computers to present adequate descriptions of molecular species.

'The future of molecular graphics: invited lecture' **Lan- gridge, R and Ferrin, T E** Computer Graphics Laboratory, University of California, CA 94143, USA

Application of the tools of such an explosively growing field, computer science, to equally rapidly developing areas such as genetic and protein engineering and drug design, make prediction unwise. We will therefore first describe the developments about which we are fairly certain, ie the direction of the UCSF Computer Graphics Laboratory over the next year or two, and devote only a little time at the end to speculation beyond that. The overall strategy of the laboratory is to combine the methods of numerical analysis, artificial intelligence and computer graphics into a system which is transparent to the user and with easily exportable software. Building on our present software (based on the Bell Laboratories UNIX operating system, which we adopted in 1976 and is now available on hosts from micros to main frames) we will install several workstations of differing levels of complexity communicating over a 10M bit/s local area network. The graphics will range from simple bitmapped black and white displays to high-performance colour vector systems. Numeric calculations for interaction geometries and energies will be done at all levels, aided by special purpose attached scientific processors. The artificial intelligence aspects will be concentrated on special-purpose LISP machines devoted to both general symbolic manipulation (eg protein chain folding) and the development of heuristics and expert systems. Supported by research grants NIH RR-1081, DOD DAAG29-83-G-0080 and a Guggenheim Fellowship.

'A low cost, high resolution colour raster graphics workstation for molecular modelling' **Lindley, M R, White, D N J and Tyler, K** Chemistry Department, The University, Glasgow G12 8QQ, Scotland

A 16-bit microcomputer using the Intel 8086 microprocessor, with 1/2 megabyte of memory and a graphics processor was used to develop a program for molecular modelling. The graphics processor is based on the Thomson EF9365 graphic processor chip, which has a resolution of 512×512 and writes 1.5 million pixels/s. A standard RGB monitor was used for the display. The system cost approximately £7500.

The program known as CHEMMOD (chemical molecular modelling system) displays molecules as colour-coded stick models. The program is user friendly, is menu driven using a lightpen and permits realtime rotation, translation, scaling and other standard manipulations of the molecules in space filling and stick formats. Space filling representations and red/green stereo views of molecules are also available for the display. Comparison of structures is provided for with a least-squares superimposition procedure. Structure building routines are included which enable