

overall dynamics are strong enough to produce chaos, lead to situations in which conventional methods of data analysis fail to reveal the density-dependent mechanisms that are built into the model (and I cited an example drawn from experimental data³). Other kinds of noise in parameters characterizing the behaviour of individuals create no such problems. Why some situations should result in difficulties in detecting density dependence, while others do not, remains the subject of investigation.

Mountford and Rothery's observation that no problems arise in their simple, phenomenological model is interesting. But it does not really address the main point in the paper by Hassell *et al.*³ or my News and Views article¹.

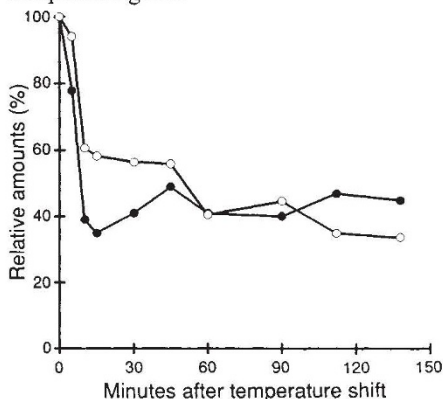
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Is yeast TCP1 a chaperonin?

SIR — G. North in his News and Views article¹ reports that the thermophilic bacterium *Sulfolobus shibatae* contains a gene, *TF55*, that is very similar in sequence to *TCP1* (ref. 2), suggestive of similar function. *TCP1* has a similar nucleotide sequence in mouse³, human³, hamster², *Drosophila melanogaster*⁴ and *Saccharomyces cerevisiae*⁵, making it a ubiquitous gene.



S. cerevisiae *TCP1* and actin transcript levels of wild-type strain M10 (ref. 5) after heat-shock. Relative transcript levels of *TCP1* (solid circles) and, for comparison, actin (open circles). Cultures were shifted at time 0 from 24 to 38 °C. The filter used to hybridize the *TCP1* probe was rehybridized with an *ACT1* probe. The absolute amounts of radiolabelled probe hybridized to specific bands was quantified on a Betascope Blot analyser (Betagen). One hundred per cent of counts for *TCP1* and actin represent 1,810 and 6,487 counts, respectively.

The data concerning the heat-shock properties of *TF55* in archaeobacteria do not extend to its homologues in yeast and *Drosophila*. In yeast, the opposite is true. Transcription of *TCP1* is repressed during heat stress as shown in the figure. Within 10 min after temperature shift from 24 to 38 °C the *TCP1* transcript level decreases by about 60%, and remains low thereafter. The upstream non-coding sequence of yeast *TCP1* does not contain the consensus elements required for heat-induced transcription⁶. In agreement with the findings in yeast, the *D. melanogaster* homologue⁴, which maps to the chromosomal location 94B, is not associated with either the major or minor heat-shock puffs⁶ and thus is not in a chromosomal region known to encode a heat-inducible protein.

Another important characteristic of most chaperonins, that they are members of a multigene family, is not shared by the *Drosophila* or yeast *TCP1* genes. The *TCP1* genes in both organisms have no detectable sequence homologues and thus do not appear to be members of a family of related genes. If yeast and *Drosophila* *TCP1* are indeed cytoplasmic chaperonins, they do not share two important attributes found to be present in most chaperonins. This would make *TCP1* a remarkable chaperonin.

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Crystallization by centrifugation

SIR — The Scientific Correspondence by J. E. Pitts¹ about the crystallization of a protein by centrifugation is not unprecedented. First beef liver catalase and later fungal (*Penicillium vitale*) catalase were crystallized by ultracentrifugation and reported in the Soviet journal *Kristallografiya*^{2,3}. The work was later published in English in *Soviet Physics, Crystallography*^{4,5}. V. Barynin presented this work at the FEBS advanced lecture course dedicated to the crystal growth of biological macromolecules (Bischenberg, Alsace, 19–25 July, 1987)⁶.

The Russian authors mentioned that the first virus coat protein was crystal-

lized in an ultracentrifuge as early as 1936⁷. Ironically, a model of the three-dimensional structure of *P. vitale* catalase, determined using the crystals grown by ultracentrifugation⁸, was built in the early 1980s on the graphics system at Birkbeck College, London.

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PITTS REPLIES — Abad-Zapatero correctly refers to the production of catalase and tobacco mosaic virus (TMV) protein crystals in an ultracentrifuge^{7,8}. In my Scientific Correspondence I alluded to this effect when I indicated that “high g-force” had been used successfully to approach the problem of crystallization. But apart from being crystallized while rotating in a centrifuge, the two methodologies are fundamentally distinct. The catalase crystals were grown at 26,000g in a high-speed ultracentrifuge over 180 hours (7½ days) in the presence of a precipitant 2-methyl-2,4-pentanediol (MPD), while the large particle size TMV crystals are produced at 40,000g^{7,8}.

The crystals of the aspartic proteinase from *Trichoderma reesei* are grown rapidly (2½ hours), at low speed and ionic strength, during rapid concentration through an ultrafiltration membrane at quite low (3,000g) centrifugal force. The nearest comparable method reported in the literature would be that of concentration dialysis⁹. The concentration process to produce protein crystals being achieved either by pressure dialysis under nitrogen¹⁰ or by vacuum dialysis through conical collodian membranes^{11,12}. The commercial availability of centric concentrators (Amicon) and of suitable slow-speed centrifuges with a fixed-angle rotor in most laboratories involved in protein purification should mean that the technique could be used more generally.

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