

regions. Amino-acid side chains exhibiting weak electron density were built with steric and geometric constraints. We generated the figures with GRASP, Molscript and Raster3D.

Structural comparisons

We carried out structural comparisons against 251 Fab and 11 $\alpha\beta$ TCR models. Superpositions were performed using LSQMAN²⁸. The inter-domain, intra-chain angles (for instance $V\gamma-C\gamma$) were calculated as the supplement of the rotation angle necessary for superposition of the conserved α -carbon atoms of the B and F strands of each domain. Because the $V\gamma-V\delta$ interface involves the A'GFCC' C' β -sheet whereas the $C\gamma-C\delta$ interface involves the ABED β -sheet (Fig. 1), V domain strands B and F were explicitly superimposed onto C domain strands F and B. The elbow angle between the V and C domains was calculated as the angle between the pseudo-dyads, described by direction cosines, which result from a superposition of V or C domains. Buried interfaces were calculated with CNS using a probe radius of 1.4 Å. Structures of $\alpha\beta$ TCRs and Fabs are referred to by their Protein Data Bank identification codes (<http://www.rcsb.org>).

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Supplementary information is available on Nature's World-Wide Web site (<http://www.nature.com>) or as paper copy from the London editorial office of Nature.

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Correspondence and requests for materials should be addressed to T.J.A. (e-mail: tallison@niaid.nih.gov) or D.N.G. (e-mail: garboczi@nih.gov). Coordinates have been deposited at the Protein Data Bank (identification code 1hxm) and will be released on publication.

corrections

Plant diversity enhances ecosystem responses to elevated CO₂ and nitrogen deposition

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Nature **410**, 809–812 (2001).

This paper should have included the following acknowledgement:

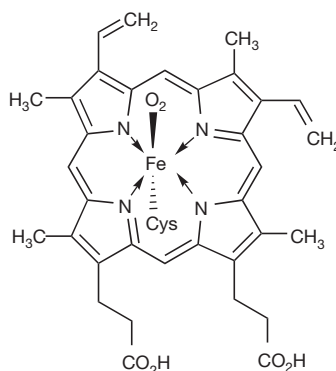
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Enabling the chemistry of life

C. Walsh

Nature **409**, 226–231 (2001).

In Fig. 4 of this *Insight* article, the P450 haem active site is wrongly shown as a uroporphyrinogen I rather than a protoporphyrin IX macrocycle. The correct version of the molecule is shown here. □



There is thus no need to invoke basal tractions due to subduction to explain pop-up structures, as has often been done for the western Americas^{27,28,29}.

The absence of slip beneath the Bhutan Himalaya in 1897 suggests that the 400-km region between the great Himalayan ruptures of 1934 and 1950 (Fig. 1) has remained a seismic gap for at least the past two centuries³⁰. At the higher end of our estimated slip rates, the faults bounding the Shillong plateau could absorb one-third of the inferred Himalayan contraction rate of 18 mm yr⁻¹ (ref. 31), correspondingly increasing the interval between great earthquakes in the Bhutan Himalaya.

Our conclusions also raise issues concerning the seismic hazard potential of the Shillong plateau. The >300-km length of the Dauki fault has not slipped recently, but were it to slip in a single earthquake its potential maximum magnitude ($M \geq 8$) would constitute a significant seismic threat to nearby densely populated regions of Bangladesh, and to the very large city of Dhaka less than 150 km to the south (Fig. 1). The interval between these giant plateau-building earthquakes fortunately exceeds 3,000 years. □

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Plant diversity enhances ecosystem responses to elevated CO₂ and nitrogen deposition

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Human actions are causing declines in plant biodiversity, increases in atmospheric CO₂ concentrations and increases in nitrogen deposition; however, the interactive effects of these factors on ecosystem processes are unknown^{1,2}. Reduced biodiversity has raised numerous concerns, including the possibility that ecosystem functioning may be affected negatively^{1–4}, which might be particularly important in the face of other global changes^{5,6}. Here we present results of a grassland field experiment in Minnesota, USA, that tests the hypothesis that plant diversity and composition influence the enhancement of biomass and carbon acquisition in ecosystems subjected to elevated atmospheric CO₂ concentrations and nitrogen deposition. The study experimentally controlled plant diversity (1, 4, 9 or 16 species), soil nitrogen (unamended versus deposition of 4 g of nitrogen per m² per yr) and atmospheric CO₂ concentrations using free-air CO₂ enrichment (ambient, 368 μmol mol⁻¹, versus elevated, 560 μmol mol⁻¹). We found that the enhanced biomass accumulation in response to elevated levels of CO₂ or nitrogen, or their combination, is less in species-poor than in species-rich assemblages.

In the twenty-first century humans will live in, manage and depend on ecosystems that are less diverse^{1,2} and subjected to higher CO₂ levels and nitrogen (N) deposition rates than in recorded human history¹. Although we are beginning to understand the individual impacts of each of these factors on terrestrial ecosystems, our understanding of their interactive effects is poor at best^{1–10}. Net primary productivity and carbon (C) input to ecosystems are usually enhanced by elevated CO₂, but this seems to be related to the extent and type of other limitations^{11–20}. Nitrogen-poor ecosystems have often shown less response to elevated CO₂ than more fertile systems^{18–25}, which is important given that, worldwide,

productivity in most terrestrial ecosystems is limited by N (ref. 26) and rates of N deposition are expected to increase in the future^{1,2,6}. In addition, it has been proposed that species-poor communities may be less responsive to elevated CO₂ or N compared with diverse communities⁷.

Our study focuses on the influence of plant diversity on the response of ecosystem functions, such as productivity, to elevated CO₂ and N supply rates. Theoretically, because the range in plant traits associated with acquisition of C and N is frequently correlated with the species or functional diversity of an ecosystem^{7,10}, ecosystem responses to elevated CO₂ and N deposition may be sensitive to variation in levels of biodiversity^{4,7,10,11,13–19}. High-diversity plant communities frequently have a greater range of plant functional traits that affect C (for example, C₃ and C₄ photosynthetic pathways) and N (for example, legumes associated with N-fixing symbionts and non-leguminous species) cycling than have species-poor communities^{7,10}, and therefore have potentially greater responsiveness of ecosystem functions to elevated CO₂ and N deposition. This can occur because diverse assemblages have a greater likelihood of containing species with strong responses to resources and strong impacts on ecosystem processes compared with species-poor assemblages (that is, a sampling effect)^{27,28}. Alternatively, this can occur because the greater range of traits extant in diverse assemblages positively affects competitive associations and interactions (such as niche complementarity and positive species interactions)^{3,7,10,28}.

Testing the influence of plant diversity on the impacts of elevated CO₂ and N to ecosystem functioning requires simultaneous manipulation of plant species composition, CO₂, and N. Although there have been glasshouse studies⁸ of species responses to elevated CO₂ in monocultures versus mixtures, and field studies of monocultures and bi-species mixtures^{9,15}, this is the first field study, to our knowledge, to test the hypothesis⁷ that plant species diversity influences ecosystem-scale biomass responses to elevated CO₂ and N levels. The BioCON experimental facility (see Methods) in Minnesota, USA, was designed expressly for the simultaneous manipulation of these three factors in experimental grassland plots under field conditions, using a well-replicated split-plot experiment comprising a full factorial combination of treatment levels in a completely randomized design.

Our study includes 296 individual plots (each 2 × 2 m) distributed among six 20-m diameter experimental areas (rings). In three elevated CO₂ rings, a free-air CO₂ enrichment system²⁹ was used during the 1998 and 1999 growing seasons to maintain the CO₂ concentration at an average of 560 μmol mol⁻¹, a concentration likely to be reached this century^{1,2,6}. Three ambient rings (368 μmol mol⁻¹ CO₂) were treated identically but without additional CO₂. All plots were planted with 1, 4, 9 or 16 perennial grassland plant species randomly chosen from 16 species in 4 functional groups (C₃ grasses, C₄ grasses, C₃ legumes, C₃ non-legume forbs). Hence high-diversity treatments incorporate greater species and functional group diversity. Native and secondary grasslands in Minnesota typically contain a mixture of these four functional types. Beginning in 1998, half the plots received addi-

tional N equivalent to 4 g N m⁻² yr⁻¹, which is comparable to high rates of N deposition observed in industrialized regions¹.

Diversity, CO₂ and N treatments had significant main effects on total biomass (Table 1, Fig. 1). Above-ground and below-ground biomass both increased markedly with increasing species diversity. Elevated CO₂ predominantly stimulated below-ground biomass, whereas enriched N largely increased above-ground biomass (Table 1). Elevated CO₂ decreased and enriched N increased plant N concentration, total plant N, soil net N mineralization and soil solution N pools (Table 1).

There was no significant interaction between CO₂ and N, and elevated levels of both resources increased total biomass by 27% on average compared with ambient plots (Fig. 1). In essence, the productivity response to elevated CO₂ was not constrained by N limitation in unfertilized plots, even those lacking N-fixing legumes (data not shown). This result differs from many earlier studies^{11,18,20}, perhaps because we added smaller amounts of N (4 g N m⁻² yr⁻¹) to represent elevated N deposition, whereas other studies (such as ref. 18) added high amounts (up to 56 g N m⁻² yr⁻¹) to mimic agricultural N addition.

Although we did not detect a CO₂ and N interaction, there were significant biomass interactions between diversity and CO₂ as well as between diversity and N (Fig. 1; and Methods), largely owing to responses of below-ground biomass, which comprised three-quarters of the total biomass. As the response to elevated CO₂ and N varied among diversity treatments, we tested their effects within diversity levels, using post-hoc comparisons of the mean responses averaged over all four harvests. In the 16-species plots, all treatments

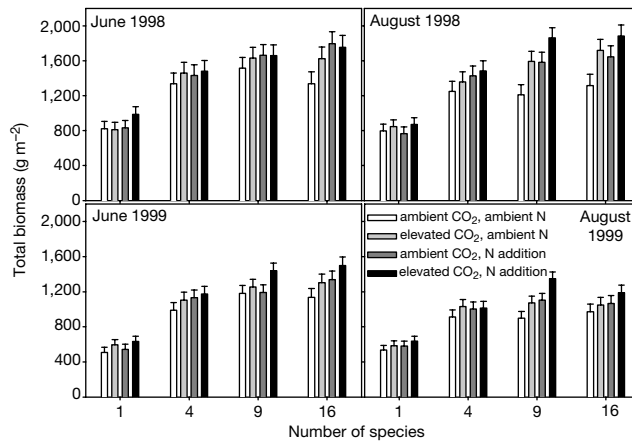


Figure 1 Total biomass (above-ground plus below-ground, 0–20 cm depth) (± 1 s.e.) for plots planted with either 1, 4, 9 or 16 species, grown at four combinations of ambient (368 $\mu\text{mol mol}^{-1}$) and elevated (560 $\mu\text{mol mol}^{-1}$) concentrations of CO₂, and ambient N and N addition (4 g N m⁻² yr⁻¹). Biomass data are shown for each of four harvests (June and August in both 1998 and 1999). The biomass (total and/or below-ground) response to elevated CO₂, enriched N, or both, differed significantly among diversity treatments at every harvest.

Table 1 Average two-year responses of experimental grassland communities to elevated atmospheric CO₂ concentrations, N deposition, and species diversity

Parameter	R ²	F ratio	CO ₂ % change	N % change	Species diversity % change
Total biomass	0.54	16.9	+12**	+13***	+98***
Above-ground biomass	0.39	9.2	+8	+23***	+81***
Below-ground biomass	0.50	15.0	+14*	+9*	+105***
Whole plant %N	0.29	5.9	-13***	+14***	-26***
Total plant N content	0.47	12.9	-2	+29***	+64***
Soil net N mineralization	0.23	4.2	-15	+64*	-68***
Soil solution N (0–20 cm)	0.21	3.9	-14	+36	-76**

R² and F ratio shown for the whole model (see Methods), for which P values were always less than 0.001. Main effects (% difference, pooled across all other treatments) and significant level (\dagger , $P < 0.1$; *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$) shown are based on measurements in all 296 plots. Species diversity % effects shown compare 16-species versus 1-species plots.

with elevated levels of either CO₂ or N had significantly greater biomass than the ambient CO₂/ambient N plots. In the 9-species plots, only the elevated CO₂/elevated N treatment had significantly greater biomass than the ambient/ambient treatment. In contrast, there was no significant effect of CO₂ or N treatment, alone or in combination, on total biomass for either the 1- or 4-species levels.

Thus, across the four harvests in 1998 and 1999, the enhancement of biomass owing to either elevated CO₂ or enriched N decreased with declining diversity. In the unamended N treatment, the average stimulation of total biomass in response to elevated CO₂ was 22% in 16-species plots, 18% in 9-species plots, 10% in 4-species plots and 7% in monoculture plots (Fig. 2). Differences in plant species diversity accounted for a fivefold difference in the impacts of CO₂ fertilization on biomass accumulation. For example, whereas elevated CO₂ increased biomass by 258 g m⁻² on average in the most diverse (16-species) plots, it increased biomass by only 47 g m⁻² on average in the monocultures (Fig. 2).

Similarly, under ambient CO₂, the average enhancement of total biomass in response to N addition ranged from 25% in 16-species plots to 18%, 11% and 2% in 9-species, 4-species and monoculture plots, respectively (Fig. 2). Thus, biomass enhancement by N enrichment was much less in low-diversity plots—N deposition increased biomass by almost 300 g m⁻² in the diverse plots and by only 15 g m⁻² in the monocultures. For plots subjected to both elevated CO₂ and enriched N, biomass increased by more than 400 g m⁻² in both the 16- and 9-species plots (+35%), but only by

roughly 100–150 g m⁻² in 4-species plots (+15%) and monocultures (+17%).

By sorting above-ground biomass to species, we found that the increase in biomass with increased diversity at each resource level was due largely to the four species that were most abundant in mixtures. The four species *Achillea*, *Bromus*, *Lupinus* and *Poa* together made up more than 80% of the above-ground biomass in 16-species plots (Table 2) and represent all three of the C₃ functional groups. Three of these species were among the four most productive species in monocultures. However, all four of the species abundant in mixtures had lower average monoculture biomass at a given resource level than the average 16-species mixtures at the same resource level (Table 2) and no one species comprised more than one-third of the above-ground biomass in the 16-species mixture at any level. Thus, consistent with a multiple-species ‘sampling effect’, there was collective dominance in 16-species mixtures by three of the more productive members of the species pool, but this dominance was moderate, and other species also contributed to the greater biomass of the high-diversity plots.

The effects of species diversity on biomass accumulation responses to elevated CO₂ and N deposition also appear to result from the combination of multiple-species sampling effects and niche complementarity or positive species interactions. The four dominant species (*Achillea*, *Bromus*, *Lupinus* and *Poa*) were responsible for much of the increase in biomass in 16-species plots at elevated resource levels (that is, elevated CO₂, enriched N, or both) and showed large responses to these treatments when grown in monoculture (Table 2; for example, *Achillea* and *Lupinus* for elevated CO₂; *Poa* and *Bromus* for elevated N).

Although the response of *Poa* monocultures to N fertilization was large enough to suggest that it may have the capability of explaining most of the increased biomass response of the 16-species mixture if heavily dominant, *Poa* was not more than one-third of the total above-ground biomass in those mixtures, indicating that other species are also important. For elevated CO₂ at ambient N, the stimulation of total biomass (+258 g m⁻²) in 16-species mixtures was greater than the stimulation in monoculture of any species to elevated CO₂. Thus, the increase in biomass of 16-species plots with increasing resource supply was substantially due to the presence and response of species that do respond strongly in monocultures (that is, interpretable as mainly a sampling effect), but they collectively increased more when together than when in monoculture (a niche effect).

In addition, different species were responsible for the enhanced responses to different combinations of CO₂ and N, again suggesting niche differentiation. The increased total biomass response of 16-species plots to elevated CO₂ at ambient N (of +22%, Figs 1 and 2) was largely due to the response of *Achillea*, *Lupinus* and *Bromus*, each of which maintained a roughly similar fraction of a higher total above-ground biomass under elevated than ambient CO₂ (Table 2). At either ambient or elevated CO₂, the enhanced biomass in N-enriched 16-species plots as compared with ambient/ambient plots was due to increases in biomass by *Poa*, *Bromus* and *Achillea*, in that

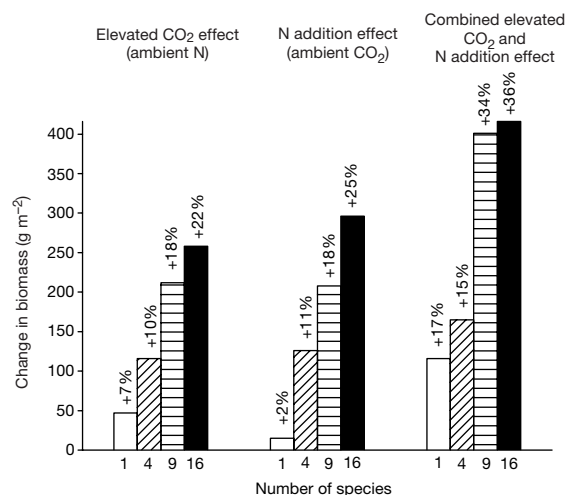


Figure 2 Change in total (above-ground plus 0–20 cm below-ground) biomass (compared with ambient levels of both CO₂ and N) in response to elevated CO₂ alone (at ambient soil N), to enriched N alone (at ambient CO₂), and to the combination of elevated CO₂ and enriched soil N, for plots containing 1, 4, 9 or 16 species. Data were averaged for 4 harvests over 2 yr. Per cent change is shown above each histogram for each diversity treatment.

Table 2 Total biomass of 16-species mixtures and of monocultures of the dominant species in the 16-species mixtures

Species	Ambient CO ₂ , ambient N		+CO ₂ , ambient N		Ambient CO ₂ , +N		+CO ₂ , +N	
	Monoculture biomass (g m ⁻²)	% of 16-species mixture	Monoculture biomass (g m ⁻²)	% of 16-species mixture	Monoculture biomass (g m ⁻²)	% of 16-species mixture	Monoculture biomass (g m ⁻²)	% of 16-species mixture
<i>Achillea</i>	1,124	34%	1,314	33%	848	31%	1,321	34%
<i>Poa</i>	979	22%	1,089	19%	1,322	33%	1,407	27%
<i>Lupinus</i>	509	15%	672	17%	449	6%	670	9%
<i>Bromus</i>	1,021	11%	989	12%	1,228	14%	1,082	14%
All other species	590	18%	615	19%	587	16%	664	16%

Totals are above-ground plus below-ground biomass, at all four combinations of CO₂ and N. Also shown is the % of total above-ground biomass for each of the dominant species in the 16-species mixtures at each CO₂ and N combination.

order, despite a decline in *Lupinus* biomass.

These results suggest that multiple-species sampling effects, niche complementarity and positive species interactions^{3,7,10,28} jointly help to explain the greater responses of diverse than species-poor plots to elevated levels of the two major global change factors: atmospheric CO₂ and N deposition. Moreover, the central finding of this study—that changes in plant diversity influence the magnitude of CO₂ and N impacts on ecosystem functioning—is important regardless of which set of mechanisms are operating.

Our study raises concerns about the consequences of widespread changes in plant composition and diversity in ecosystems worldwide for responses to other global changes^{1,2}, because the results show that, in response to elevated levels of CO₂ and N, ecosystems with decreased diversity may acquire less C and biomass than ecosystems with greater diversity. It is unclear whether the responses observed in 1998–1999 will change with time as the composition of the communities changes, the climate varies, and long-term soil microbial feedbacks occur^{20–25}, or whether natural or managed vegetation would respond similarly to combinations of diversity, CO₂ and N. Nonetheless, our results suggest that the reduction of diversity occurring globally may reduce the capacity of ecosystems to capture additional C under conditions of rising atmospheric CO₂ concentrations and N deposition levels. □

Methods

BioCON experiment

The BioCON (Biodiversity, CO₂ and N) experiment (<http://swan.lter.umn.edu/biocon/>) is located at the Cedar Creek Natural History area in Minnesota, USA. Plots were established on a secondary successional grassland on a sandy outwash soil after removing the previous vegetation. The experimental treatments were arranged in complete factorial combination of CO₂ (ambient or 560 μmol mol⁻¹), species number (1, 4, 9, and 16) and N level (control and fertilized). The species numbers were chosen as the squares of 1, 2, 3 and 4 to represent roughly equal effective differences in diversity, on the basis of earlier studies⁴. Each plot was planted in 1997 with 12 g m⁻² of seed partitioned equally among all species planted in a plot.

The design consisted of a split-plot arrangement of treatments in a completely randomized design. CO₂ treatment is the whole-plot factor and is replicated three times among the six rings. The subplot factors of species number and N treatment were assigned randomly and replicated in individual plots among the six rings. For each of the four combinations of CO₂ and N levels, pooled across all rings, there were 32 randomly assigned replicates for the plots planted to 1 species, 15 for those planted to 4 species, 15 for 9 species, and 12 for 16 species. Beginning in 1998, the plots assigned to the N addition treatment were amended with 4 g N m⁻² yr⁻¹, applied over three dates each year. CO₂ was added in elevated treatments during all daylight hours from 9 April to 16 October 1998, and from 20 April to 9 November 1999. Although there was modest variation in CO₂ concentrations spatially within and across rings, the average CO₂ concentrations were not more than 1–2 μmol mol⁻¹ different among rings, or among diversity or N treatment levels averaged within and across rings (see Supplementary Information).

Species and biomass measurements

The 16 species used in this study were all native or naturalized to the Cedar Creek Natural History Area. They include four C₄ grasses (*Andropogon gerardii*, *Bouteloua gracilis*, *Schizachyrium scoparium*, *Sorghastrum nutans*), four C₃ grasses (*Agropyron repens*, *Bromus inermis*, *Koeleria cristata*, *Poa pratensis*), four N-fixing legumes (*Amorpha canescens*, *Lespedeza capitata*, *Lupinus perennis*, *Petalostemum villosum*) and four non-N-fixing herbaceous species (*Achillea millefolium*, *Anemone cylindrica*, *Asclepias tuberosa*, *Solidago rigida*), and all are referred to by genus elsewhere. Monocultures of all species were replicated twice at all CO₂ and N levels. The 4- and 9-species plots were random selections from all species. Plots were regularly weeded to remove unwanted species. In June and August of each year, we assessed above- and below-ground (0–20 cm) biomass, plant C and N, and soil N (see Supplementary Information). Soil net N mineralization rates were measured once each year. Above-ground biomass was sorted to species at each harvest. The species richness of clipped above-ground biomass samples was 1.0, 3.8, 8.0 and 13.6 species on average for plots planted with 1, 4, 9 and 16 species, respectively, on the basis of the number of species identified in each plot from the four above-ground harvests (each 0.1 m²).

Statistical analysis

In analysis of variance all treatment effects were considered fixed. Using *F*-tests, the effect of CO₂ (1 degree of freedom, d.f.) was tested against the random effect of ring nested within CO₂ (4 d.f.). The main effects of species number (3 d.f.) and N (1 d.f.), and interactions between CO₂ and N were tested against the residual error. The main effect of species number and its first-order interaction terms were partitioned into single-degree-of-freedom contrasts for linear, quadratic and cubic terms to test for interactions between diversity and either CO₂ or N treatments, and additionally to test hypotheses about

predetermined contrasts of elevated resource levels (elevated CO₂, enriched N, or both) versus the ambient/ambient conditions. There were interactions for below-ground biomass between species diversity and CO₂ treatments in the August 1998 (*P* < 0.05), June 1999 (*P* < 0.10) and August 1999 (*P* < 0.10) harvests, and between species diversity and N treatments in the June 1999 (*P* < 0.05) and August 1999 (*P* < 0.05) harvests. For the pre-planned contrast between the ambient/ambient and the elevated CO₂/enriched N treatments, there were significantly different responses (for below-ground and total biomass) for different diversity treatments at the June 1998 (*P* < 0.05), August 1998 (*P* < 0.05) and June 1999 (*P* < 0.05) harvests. Given interactions between species diversity and resource treatments, post-hoc Student's *t*-tests were also conducted to compare individual resource treatments within species diversity levels. All analyses were conducted for each harvest, for each year (pooling harvest data by plot), and across years averaged (pooling data for all harvests by plot), with similar results, although there was harvest-to-harvest variation (see Fig. 1).

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Enabling the chemistry of life

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Enzymes are the subset of proteins that catalyse the chemistry of life, transforming both macromolecular substrates and small molecules. The precise three-dimensional architecture of enzymes permits almost unerring selectivity in physical and chemical steps to impose remarkable rate accelerations and specificity in product-determining reactions. Many enzymes are members of families that carry out related chemical transformations and offer opportunities for directed *in vitro* evolution, to tailor catalytic properties to particular functions.

The myriad chemical transformations carried out by every living organism are enabled by hundreds to thousands of proteins (enzymes) and, less frequently, RNAs (ribozymes), which have catalytic activity for conversion of a particular set of substrates to specific products. Some of these reactions are carried out by related families of protein biocatalysts, which act generically in the same way but exert specific recognition for transformation of a particular substrate molecule. For example, the orderly control of the location and lifetime of proteins in cells is managed by dozens of related proteases that hydrolyse peptide bonds of protein substrates in ways that are controlled in time and space. Proteases can be exquisitely specific for a particular peptide bond in a protein substrate, or they can be relentlessly nonspecific: the former set of proteases are involved in turning on biological signals, the latter in the clean-up phases of degradation and protein turnover.

When cells respond to external messenger molecules, such as the protein growth factors and hormones erythropoietin and insulin, or small-molecule hormones such as adrenaline or prostaglandins, signalling pathways are set in motion by catalytic action of cascades of protein kinases. The protein kinases are built from a small set of architectural types, and all catalyse phosphoryl transfer from ATP to the side-chain hydroxyl of serine, threonine or tyrosine residues. There are hundreds of such kinases in animal genomes. Selectivity is imposed on this generic chemical phosphorylation reaction by protein–protein interactions between a given kinase and its protein substrate and by cascades of such kinase/protein substrate pairs that ultimately lead to changes in activity and location of proteins, and to selective gene activation.

In addition to the large number of enzymes that act on macromolecular protein substrates, there are also enzymes that engage in truly sophisticated chemistry on small

Figure 1 Diverse chemical reactions facilitated by biocatalysts. ACC lyase, 1-aminocyclopropane-1-carboxylate lyase; IPNS, isopenicillin *N* synthase; OMP decarboxylase, orotidine-5'-phosphate decarboxylase.

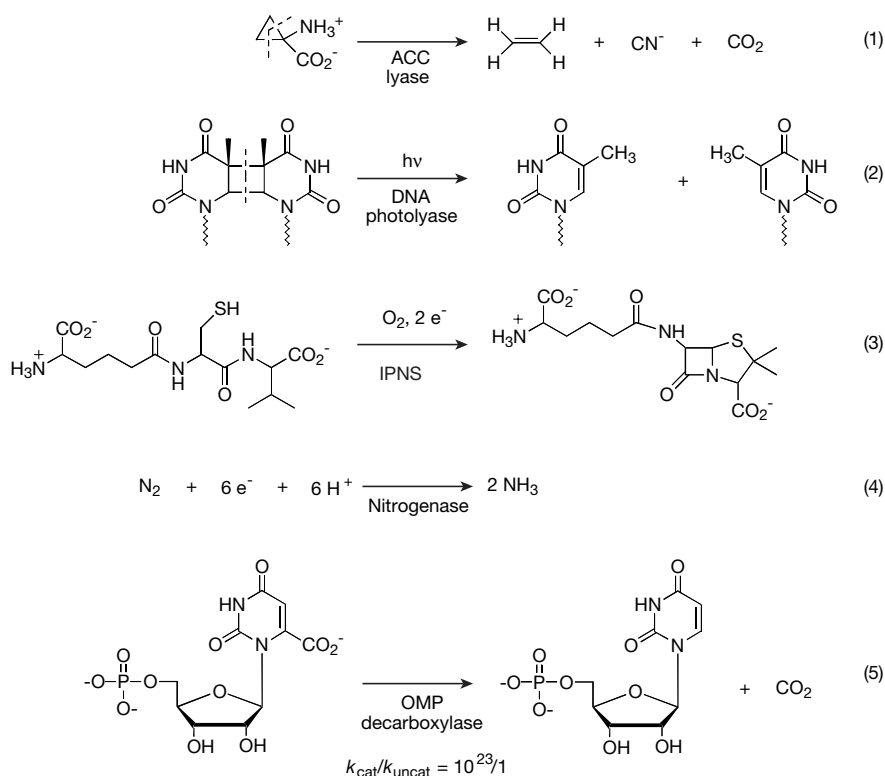
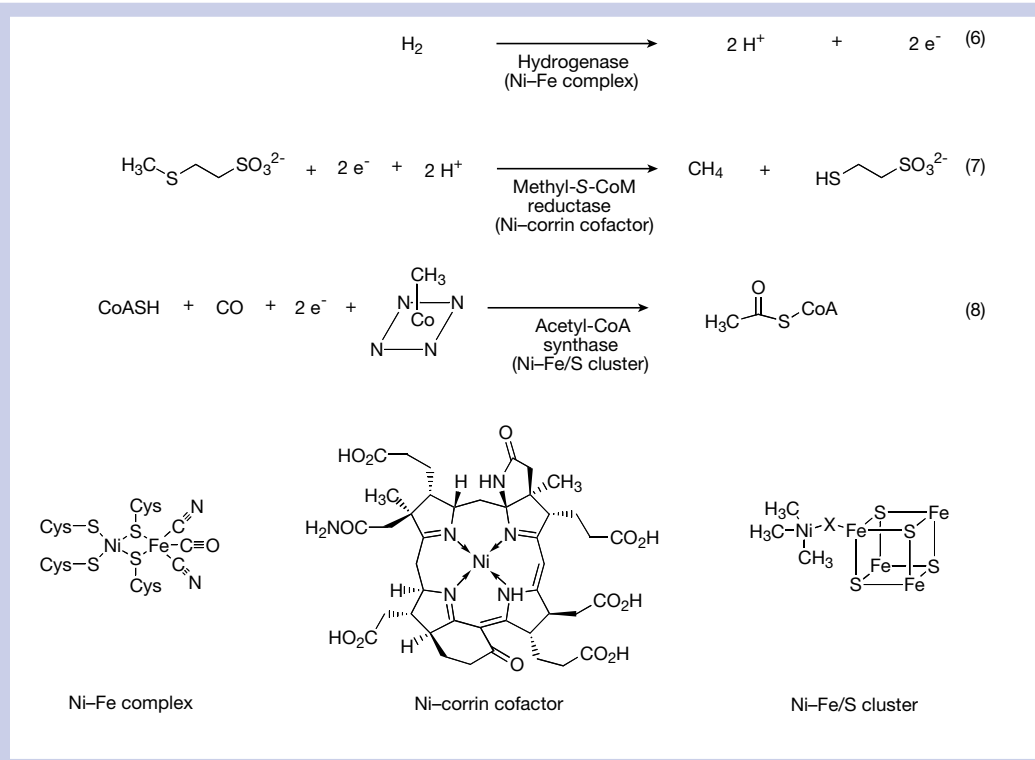


Figure 2 Nickel-based enzymatic transformations in methanogenic archaeobacteria.



organic molecules. The fragmentation of 1-aminocyclopropane-1-carboxylate to the fruit-ripening hormone ethylene¹, the photon-induced 2 + 2 cycloreversion of thymine dimers to repair DNA damaged by ultraviolet light², the bis-cyclization of the tripeptide aminoacyl-cysteine-D-valine (ACV) to isopenicillin N (ref. 3), and the reduction of dinitrogen (N₂) to two molecules of ammonia (NH₃) during nitrogen fixation⁴ are just a few examples of the range of biological chemistry facilitated by biocatalysts (Fig. 1). Enzymes as biocatalysts are remarkable not only in themselves, but also for the inspiration and guidance they provide to synthetic organic and inorganic chemists striving to reproduce and expand nature's chemical repertoire. Several of the useful attributes of biocatalysts, such as their use as reagents for chemical synthesis and scale-up, and directed evolution to tailor chemical transformations, are explored in other articles in this Insight.

Biocatalysts and their *ex vivo* utility

Biocatalysts carry out the chemistry of life, the controlled chemical transformations in primary metabolism and the generation of natural-product diversity in secondary metabolism of plants and microbes. Classically, the subset of proteins with catalytic activity — the enzymes — has been the focus of biocatalysis research. But there is an increasing focus on catalytic RNA (ribozymes), the discovery of which in the 1980s supported the arguments for an 'RNA world'^{5,6} antecedent to the contemporary world where proteins are the workhorse biocatalysts. Most recently, Joyce and co-workers⁷ have reported catalytic DNA molecules, and directed evolution of both RNA and DNA biocatalysts will continue to expand their potential. The current set of RNA and DNA catalysts have been assayed and developed for activities in nucleic-acid replication and in protein synthesis^{8,9}, but it remains to be seen how suitable they will be for the chemically diverse reactions encompassed by existing enzyme catalysts.

The twin hallmarks of enzyme biocatalysts are the remarkable specificities and sometimes phenomenal rate accelerations achieved. A typical enzyme, with a relative molecular mass of 50,000 (*M_r* 50K), is comprised of 450 amino-acid residues: 19 chiral L-amino acids and glycine. If glycine makes up 10% of the residues, then there are at least 400 residues with chiral centres to provide an asymmetric microenvironment for substrate binding and subsequent chemical

transformation in the enzyme's active site. This is the underlying structural basis for the action of all enzymes as chemoselective and regio- and stereospecific catalysts. In terms of rate accelerations, the relative values over nonenzymatic rates of transformation are often 10¹⁰, for example for protease-mediated hydrolysis of peptide bonds, and can reach 10²³ in the example of orotidine decarboxylase in the pyrimidine biosynthetic pathway¹⁰ (reaction 5 in Fig. 1). In absolute terms, enzymes have turnover numbers from as slow as one catalytic event per minute to 10⁵ per second (as in the hydration of CO₂ to HCO₃⁻ by carbonic anhydrase)¹¹.

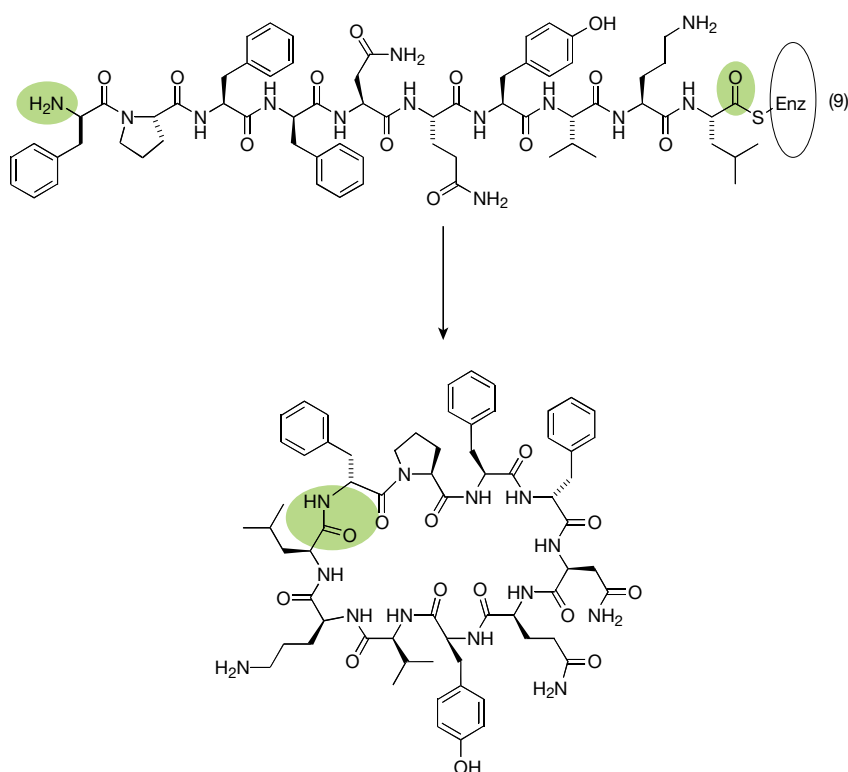
These two attributes of enzymatic biocatalysts have spurred much investigation into both the structural and mechanistic bases of the chemical transformations and have stimulated much of the study of enzymes in chemical synthesis (see review in this issue by Koeller and Wong, pages 232–240). *In vivo*, enzymes operate in buffered aqueous environments with ionic strength and pH control, although microbes that live at extremes of temperature and pH are of particular current interest because of the stability of their constituent enzymes. Much attention in biocatalyst process design (see accompanying review by Witholt *et al.*, pages 258–268) is on how to prolong useful lifetimes of enzyme catalysts and to have them operate in media not ordinarily compatible with life.

The past two decades have also witnessed an intense exploration of catalytic antibodies¹². To prepare these antibodies, ligands are synthesized that typically mimic transition states of particular chemical transformations, such as ester hydrolysis, amide synthetase and Claisen condensation. Monoclonal antibodies are then selected that display high-affinity binding to the ligands, thus enriching for antibody proteins with a binding-site geometry complementary to the shape of the true transition state. Some of the antibodies selected in this way show catalysis of the desired reactions, with the selectivity and rate accelerations expected for chiral protein-based catalysts^{13,14}. But low catalytic turnover numbers have so far limited the use of catalytic antibodies in chemical synthesis or process work.

Biocatalysts or biomimetic catalysts?

With their unerring stereoselectivity and high catalytic efficiency, nature's enzymatic catalysts have been a stimulus and counterpoint

Figure 3 Cyclization catalysed by the thioesterase domain of tyrocidine synthetase.



for generations of chemists who have designed and tested bioorganic and bioinorganic versions of biomimetic catalysts, whether for example to mimic macrocyclizations of natural products or to produce analogues of hydrogenase or nitrogenase catalysts or the photosynthetic splitting of water¹⁵. The mimics may operate under harsher solvent and temperature conditions, and may be more robust in terms of lifetime (if not throughput per catalyst molecule). When organic coenzymes (such as flavins, pyridoxal or thiamin) or inorganic cofactors (iron/sulphur clusters, metalloporphyrins) are crucial components of the enzymatic catalysis, the biomimetic and natural catalysts often show design convergence and may recapitulate some of the steps in biocatalyst evolution. The three nickel enzymes in methanogenic bacteria (thought to be contemporary descendants of primordial organisms), which carry out nickel-based hydrogenation, nickel-based methyl thioether reduction to methane, and nickel-based carbonylation of a methyl co-substrate to produce acetate, can be viewed as such an intersection^{16,17} (Fig. 2).

When is it worthwhile for the synthetic or process chemists to reject synthetic reagents and catalysts in favour of enzymes to carry out a specific transformation? This may vary with individual preference and each case must be judged on its own merits. Lipases and other hydrolases have clear advantages in kinetic resolutions of intermediates (see below), penicillin acylases have long been a mainstay of semisynthetic processes in the β -lactam antibiotic industry, and enzymatic aldol condensations have shown their worth in complex oligosaccharide syntheses¹⁸.

Chemical transformations well suited to enzymes

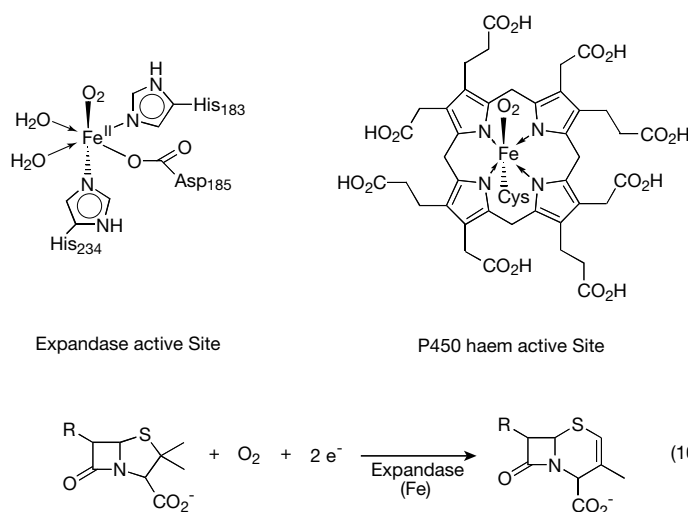
The accompanying review by Khosla and Harbury (pages 247–252) explores the multimodular enzymes that function as molecular solid-state assembly lines for the generation of thousands of polyketide natural products and non-ribosomal peptide antibiotics, including important medicinal compounds such as erythromycin, rapamycin, epothilone, lovastatin, penicillins, cyclosporin and vancomycin^{19–21}. These sequentially elongating acyl transfers seem

particularly apt loci for use as enzymatic rather than biomimetic catalysis. Some of the assembly lines, such as those for erythromycin or cyclosporin, produce the intramolecularly cyclized macrolactones or macrolactams. It has recently been shown²² that the last 30K (thioesterase) domain of the 724K protein assembly line of tyrocidine synthetase retains the ability to cyclize 9–11-residue peptidyl thioesters with regio- and stereoselectivity, raising the prospect for practical enzymatic macrocyclizations by a robust, small protein fragment (Fig. 3, reaction 9).

The reprogramming of the component enzyme domains of these assembly lines to create new, unnatural ‘natural’ products is one of the goals of combinatorial biosynthesis. The order of the enzymatic domains in the assembly lines specifies which monomer substrates are activated, condensed and elongated. So altering the order and permutations of these domains offers the chance to control product structure. The directed evolution of the catalytic domains of polyketide synthase (PKS) and non-ribosomal peptide synthetase (NRPS) assembly lines by gene shuffling and other approaches (see accompanying review by Arnold, pages 253–257) can create designed diversity in complex natural products.

Once the nascent products have been released from the PKS and NRPS assembly lines, the polyketide or polypeptide may require further enzymatic transformations to attain antibiotic properties. This is the case for penicillins, vancomycin and erythromycin, to cite just three important examples¹⁹. Baldwin and co-workers^{23,24} showed that the tripeptide ACV is oxidatively transformed to the 4–5 bicyclic β -lactam ring system by isopenicillin *N* synthase (IPNS; Fig. 1, reaction 3). IPNS is a member of a substantial family of iron-containing enzymes that use Fe^{2+} to activate both O_2 and the specific co-substrate for complex redox chemistry²⁵. In IPNS, both atoms of dioxygen are reduced to water and the ACV tripeptide undergoes four-electron oxidation and directed C–S bond and C–C bond formation as the β -lactam forms. A cousin of IPNS, the expandase enzyme, is used by cephalosporin-producing organisms to expand the five-membered ring in penicillins to the six-membered ring in cephalosporin

Figure 4 Comparison of expandase active site with a typical haemprotein oxygenase.



antibiotics (Fig. 4, reaction 10). The ligand set around the active-site iron — one Glu, two His residues — is the same, but the reaction flux is distinct (Fig. 4). Other members of this non-haem dioxygenase family include the enzyme responsible for hydroxylating prolyl residues in procollagen to predispose it to triple-helix formation in mature collagen, the most abundant protein in the human body. There are clear potential benefits to understanding the molecular basis for how the high-valent oxo-iron reagents are controlled and directed to flawlessly different chemical outcomes in the members of this redox enzyme family, so that they might be subjected to *in vitro* evolution to generate new reaction fluxes.

Many natural products, from morphine and codeine to vancomycin, undergo oxidative cyclization reactions that are regio- and stereospecific and seem to be mediated by a different superfamily of iron-containing oxidases, the cytochromes P450, with Fe²⁺ embedded in a haem macrocycle (Fig. 4). Protein superfamilies are groups of proteins with distinct chemical functions, amino-acid sequences of recognizable but sometimes marginal homology, and convergent three-dimensional structures. In the vancomycin family of glycopeptide antibiotics there are three crosslinks that convert an acyclic heptapeptide, the product of the NRPS assembly line, into a rigid scaffold, crosslinked at Tyr₂-PheGly₄-Tyr₆ and PheGly₅-dihydroxyPheGly₇ (Fig. 5, reaction 11). There are three P450 cytochromes in the biosynthetic gene cluster; each might enact a regiospecific phenolic crosslink. Harnessing such catalysts for related transformations might lead to new vancomycins.

Several natural products contain tandem five-membered-ring heterocycles (oxazoles and thiazoles) that arise from enzymatic cyclization of serine or cysteine residues in peptide precursors²⁶. These include the *Escherichia coli* antibiotic microcin B17, which kills neighbouring bacteria by poisoning the enzyme DNA gyrase and thus blocking DNA replication, in much the same way as does the best-selling antibiotic ciprofloxacin²⁷ (Fig. 6). Such heterocycles are also found in the iron-chelating siderophores that act as virulence factors in infections by *Pseudomonas aeruginosa*, *Vibrio cholerae* and the causative agent of the black plague, *Yersinia pestis*^{28,29}. Enzymes that heterocyclize serine, threonine and cysteine side chains in peptides (Fig. 6, reactions 12, 13) may create either DNA-seeking or iron-chelating sites in any peptide library that could then be screened for biological activity.

Superfamilies, genomics and enzyme evolution

The iron-containing dioxygenases that include IPNS and expandase, and the cytochrome P450 variants that introduce crosslinks, comprise redox enzyme superfamilies that are good candidates for

engineering for altered catalytic properties and specificities. Genomic and proteomic searches can identify many enzyme superfamily members through amino-acid sequence homologies, in which scaffolding and structural architecture will be predictable. Some of these proteins are of unknown ('orphan') function, and the assignment of function is one of the major postgenomic challenges of proteomic research. Recent cases in the crotonase superfamily (Fig. 7, reactions 14–16) and enolase superfamily (Fig. 7, reactions 17–19)^{30–32} indicate that the active sites all generate carbanionic transition states from bound substrates and then use carbanion chemistry for directed fluxes and distinct product outcomes. These families should be fruitful starting points for directed enzyme evolution to elicit new fluxes, based on the knowledge that carbanion chemistry will be facilitated

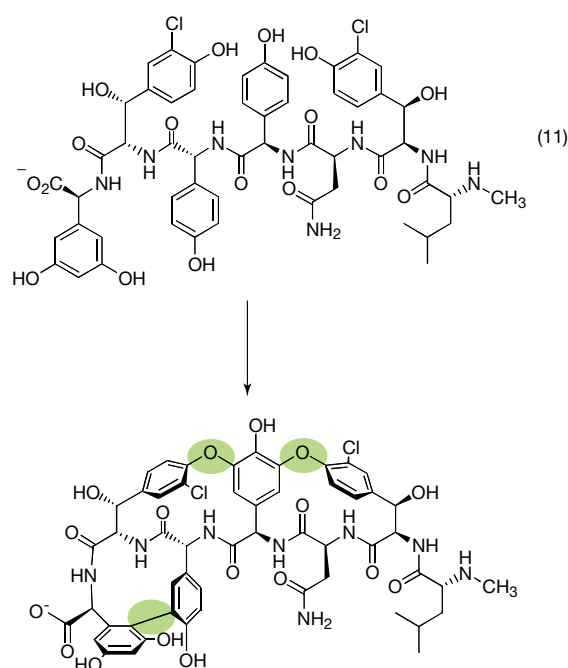
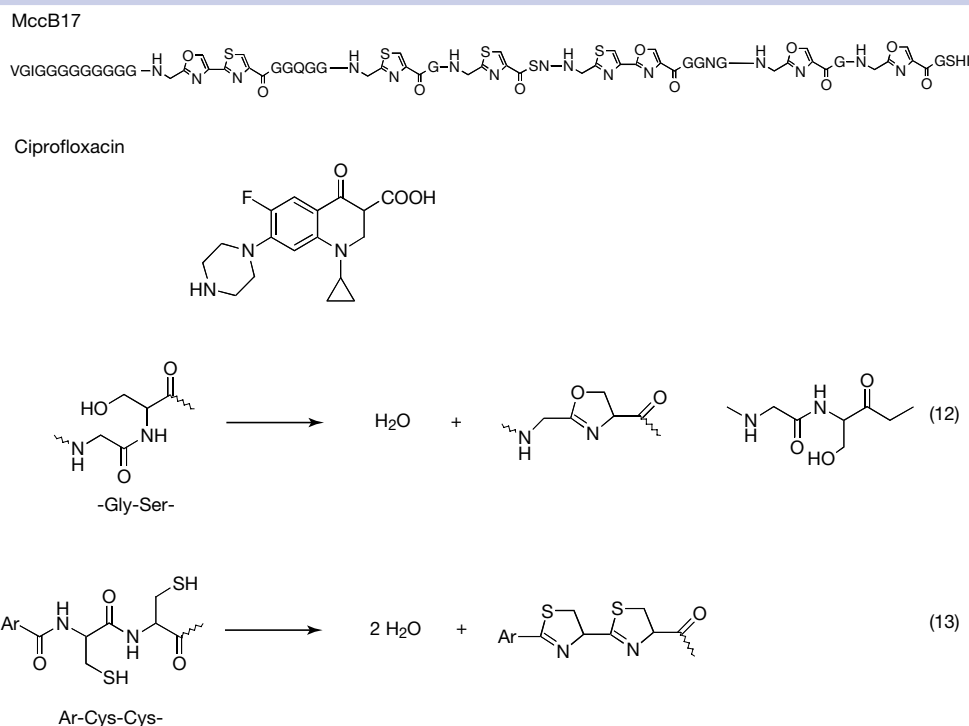


Figure 5 Crosslinking by cytochrome P450 enzymes to produce the vancomycin aglycone.

Figure 6 DNA gyrase inhibitors and biosynthesis of peptide heterocycles.



in one of the co-substrates and that binding sites can be re-engineered for electrophilic substrate components.

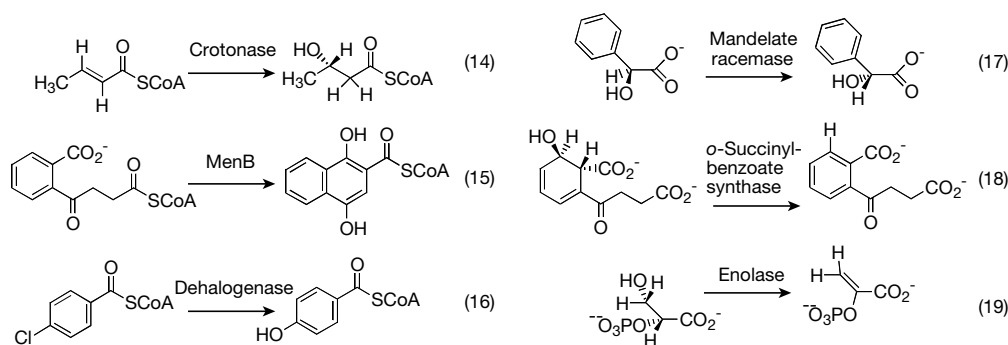
Once an enzyme has been evolved to have a detectable and desirable new activity³³, additional rounds of *in vitro* evolution can improve its stability and robustness. The biological selection methods are sufficiently powerful that one can find outcomes that are very rare biologically in a short space of time. A good example is the recent report³⁴ of expression of a functional carotenoid biosynthetic pathway in *E. coli* by selecting for bacteria that become red. The continuing progress in biological production of polyhydroxyalkanoate polymers with controlled sizes and properties³⁵ by engineering the respective polymerases increases the likelihood of economically viable production of these biodegradable plastics by biocatalysis.

Enzymes in bioremediation

One of the most active areas of applied enzymology in the past two decades has been the study of enzymes capable of bioremediation: the breakdown of organic and inorganic pollutants. There are now substantial databases of enzymes and the bioremedial transfor-

mations³⁶ they catalyse, which include the breakdown of aromatic and heteroaromatic pollutants by oxidative, reductive and hydrolytic transformations. Iron-containing dioxygenases and monooxygenases, with overlapping regio- and chemospecificities, are superfamilies that represent good starting points for application of many of the strategies noted here and in the specific accompanying articles for directed enzyme evolution to broaden substrate recognition. It is likely that bioremediation scenarios in the field will require the tailored enzymes to work in their host microbial cells rather than as *ex vivo* catalysts. Engineering of multistep metabolic pathways by introducing heterologous genes³⁷ and *in vivo* expression may well be required for efficient degradation of non-biogenic compounds. As many waste sites have a witches' brew of foreign compounds, multiple pathways engineered stably into a microbe or, more probably, mixed bacterial communities that can coexist stably, will be required. The enzymology of processing of toxic inorganic ions has also progressed in recent years to include mercury, copper, cadmium, silver, arsenic and cobalt. This might ultimately make remediation schemes for inorganic pollutants feasible³⁸.

Figure 7 Representative reactions catalysed by the crotonase superfamily and the enolase superfamily.



Conclusions

As structural genomics continues to reveal the folds and scaffolds of several members of all the principal superfamilies of enzymes, the molecular bases of recognition of substrates and directed fluxes through specific transition states to particular subsets of products will become increasingly clarified. In turn this will aid in enzyme evolution to select and detect new activities and then to incorporate improved catalytic efficiency, attributes of specificity, and structural features optimized to a given operating microenvironment. For both *in vitro* applications for a specific synthetic chemical step and for *in vivo* construction of new metabolic pathways, the applications for enzymes in practical biocatalysis will continue to burgeon. Small-molecule chemical transformation catalysed by enzymes from microorganisms that live in unusual environments or conduct chemical warfare against their neighbours have been and are likely to remain good hunting grounds for new enzyme transformations. □

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