

NEW EMBO MEMBERS' REVIEW

The black cat/white cat principle of signal integration in bacterial promoters

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Introduction

'Black cat, white cat: whatever catches the mouse is a good cat' (Chinese proverb).

Throughout the 3000 million years that bacteria have been on planet Earth, they have evolved amazing mechanisms for rapid adaptation to every imaginable type of environmental change. To give one extreme example, as little as 50 years after the introduction of the first antibiotics, very few of these drugs remain effective today at combating microbial infections (Davies, 1997). When bacteria are faced with antimicrobials, the only challenge they must overcome is to defeat the toxic effect of the compound through modification, cleavage or pumping the drug out of the cells; these processes generally involve one or just a few proteins. A far more complex environmental threat is that posed by organic chemicals, which have been discharged into many ecosystems through industrial and urban activities. Many of these chemicals are xenobiotic compounds (literally, alien to life), which include types of chemical ligatures (typically covalent C–Cl bonds) that have never been present in significant amounts in the biosphere and against which the housekeeping metabolic pathways of most microbes are generally useless (van der Meer *et al.*, 1992). Unlike antibiotics, the challenge in this case is the construction of entire biodegradation pathways that endow bacteria with the ability to grow on these otherwise unpalatable chemicals. Such an outcome involves not just one protein, but sometimes dozens, which must adapt to entirely new substrates and intermediates. However, difficult as this might be, the successful assembly of a degradation pathway does not guarantee *per se* the survival of a particular strain. Bacteria that colonize polluted sites are subject to extremely tough competition from other microbial residents of the same niche. Transcriptional regulation of biodegradative genes and operons thus becomes a critical asset for the success of a newly assembled pathway to ensure its expression at only the right moment with a minimal waste of energy (de Lorenzo and Pérez-Martín, 1996). But how do bacteria learn to respond optimally to novel environmental signals and substrates?

Most of the known functional characteristics of prokaryotic promoters come from studies that use *Escherichia coli* as the test organism. Although far more complex than *Buchnera* (one of the simplest bacteria known so far; Shigenobu *et al.*, 2000), the life cycle and the natural niches of *E. coli* can be relatively simple compared with those of not-so-distant relatives, such as pseudomonads, which thrive in soils polluted with toxic chemicals. Such rapidly adapting bacteria have become the experimental systems of choice in understanding how genes and pathways end up with regulated expression. In fact, since bacteria have been exposed to some such pollutants for only a few decades, it is possible, as discussed below, to find intermediate steps in the evolution process. In this review we summarize the features of the regulation of catabolic pathways for recalcitrant aromatic compounds that can help us to understand such a process. The conclusion is what we refer to as the 'black cat/white cat principle', which states that any regulatory mechanism is equally efficient provided that it ensures both a responsiveness to the new substrate and a suitable connection with the physiological state of the bacteria. A number of well studied cases to substantiate this notion are discussed below.

Transcriptional noise: promoters learning to respond to novel chemicals

The regulation of pathways for biodegradation of recalcitrant compounds by Gram-negative soil bacteria (mostly *Pseudomonas*, *Alcaligenes*, *Bulkholderia* and *Acinetobacter*) reveal some interesting mechanistic features by which operons acquire conditional promoters (Díaz and Prieto, 2000). The functioning of a new route depends on two major requirements that bacteria must attain to utilize the evolutionary advantage offered by the presence of fresh chemical species as potential carbon sources. One is the complement of genes encoding the whole suite of enzymes that build a pathway of reactions leading to metabolism of the compound to carbon dioxide and water. Operons destined for work in polluted sites need, in addition, an efficient transcriptional control system (de Lorenzo and Pérez-Martín, 1996). Regulated promoters are the key element that permit catabolic operons to be transcribed only when required and at levels adequate to guarantee a satisfactory metabolic return from the substrate. These two steps (assembly of a catabolic operon and acquisition of a substrate-responsive promoter) seem to be independent events, governed by different rules that operate at various times.

In general, isoenzymes that catalyse similar steps within a pathway tend to be alike at the sequence (DNA, protein) level also, even in cases where the initial substrates of the pathway are very different. This allows us to trace the

Table I. Levels of regulatory noise in catabolic promoters

Phenomenon	Examples	Reference
Minor/major gratuitous induction	Induction of the <i>upper</i> TOL pathway by <i>o</i> -xylene Induction of the lower NAH pathway by anthranilic acid Induction of the <i>alk</i> pathway by DCPK Induction of the lower TOL pathway by 2Br-benzoate Residual induction of σ 54-dependent promoters by large collections of non-substrates	Abril <i>et al.</i> (1989) Cebolla <i>et al.</i> (1997) van Beilen <i>et al.</i> (1994) Ramos <i>et al.</i> (1986) Abril <i>et al.</i> (1989); Garmendia and de Lorenzo (2000); Kahng <i>et al.</i> (2000); Jaspers <i>et al.</i> (2000)
Cross-talk between akin regulators	Benzoate-responsive activation of the <i>Pm</i> promoter of the lower TOL pathway in the absence of XylS Activation of the <i>tfd</i> gene cluster of <i>Ralstonia eutropha</i> JMP134 by TfdR in the absence of TfdT Activation of <i>clc</i> genes by CatR in the absence of cognate regulator ClcR in <i>P.putida</i> AC27 Cross-regulation of toluene monooxygenases by TbmR and TbuT in <i>Ralstonia pickettii</i> PKO1	Leveau and van der Meer (1996) Parsek <i>et al.</i> (1994) Leahy <i>et al.</i> (1997)
Regulatory takeover between related regulators	Phenol-dependent activation of <i>upper</i> TOL operon by DmpR and toluene-dependent activation of the <i>dmp</i> operon by XylR Activation of the <i>pheBA</i> promoter of <i>P.putida</i> PaW85 by CatR	Fernández <i>et al.</i> (1994) Parsek <i>et al.</i> (1995)
Promiscuous activation	Activation from solution of σ 54-dependent promoters by various regulators of the NtrC family	Pérez-Martín and de Lorenzo (1995)

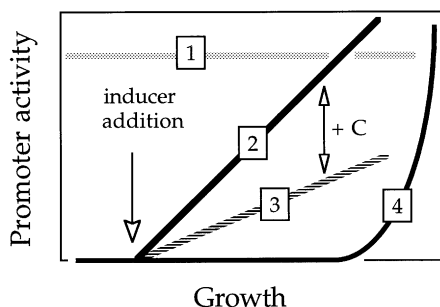


Fig. 1. Expression profiles and development of promoters responding to novel environmental signals (e.g. latest carbon sources). Expression of genes and gene clusters encoding new catabolic abilities towards an evolutionarily recent substrate may be simply achieved through constitutive expression (type 1). This may evolve further into inducible expression (type 2). The presence of other nutrients in the medium that are easier to metabolize can later influence such induction and cause a C-source inhibition (type 3). Finally, both the presence of other nutrients, growth rate, and other environmental and physiological signals can be integrated for downregulation of the promoter during rapid growth, causing the so-called exponential silencing (type 4).

origin of novel pathways to the patchwork assembly of pre-existing DNA segments bearing gene variants active on the novel substrates (van der Meer *et al.*, 1992). Excellent examples of this include the adaptation of the entire set of catabolic genes of *Pseudomonas putida* for degradation of benzoate (involving the *ben* and *cat* genes) [Cl-benzoates (Parsek *et al.*, 1994) or methyl-benzoates (van der Meer *et al.*, 1992; Ramos *et al.*, 1997)] or the recruitment of at least three catabolic segments for degradation of the very recalcitrant compound pentachlorophenol (Copley, 2000). There are no rules, however, to anticipate the type of regulator that may appear to respond to a novel chemical structure. New pathways generally start with low-level constitutive expression, on top of which increasingly specific promoters might develop. In contrast to the assembly of the enzymatic pathways, there is no specific requirement for a given type of regulator; on the contrary,

it is common to find nearly identical catabolic operons preceded by entirely different regulatory devices and proteins (de Lorenzo and Pérez-Martín, 1996). The bottom line is that a novel specificity may evolve through the leakiness of an earlier transcriptional control scheme (i.e. transcriptional noise). Any new control system should therefore start by recruiting the residual responsiveness of an already existing promoter/regulator to another signal, and then evolve by selecting changes such that residual responsiveness becomes predominant. As a consequence, the process from constitutive expression to response to a single inducer involves intermediate steps with various degrees of specificity, which can be found as remnants of the process in many catabolic pathways. Table I shows a number of cases that have been examined in detail. They include gratuitous induction (responsiveness to non-metabolizable compounds), cross-talk between structurally, but not functionally, related regulators, full replacement of one factor by another, or promiscuous activation among regulators of the same family. Pathways found in bacteria able to degrade substrates only very recently found in the environment (e.g. pentachlorophenol, polychlorobiphenyls, hexachlorocyclohexane, nitroaromatics, etc.) are often poorly regulated (Mouz *et al.*, 1999; Copley, 2000; Watanabe *et al.*, 2000), probably reflecting only an early step in the optimization of the corresponding metabolic route.

Physiological control of transcription: the phenomenon

Bacterial regulation of catabolic pathways in the environment implies not just the ability to respond to a substrate, but also whether or not expression of the whole complement of enzymes is beneficial or detrimental to ecological performance. Bacteria thriving in a polluted niche receive a range of physical and chemical signals, other than just the presence of a substrate, which need to be processed to achieve a positive or negative outcome for each specific

Table II. Biodegradative pathways subject to physiological control in pseudomonads

Pathway/operon and phenotype	Substrate	Reference
Degradation of the hydrocarbon by <i>P.putida</i> CA3 is inhibited by glutamate and citrate, not by glucose.	styrene	O'Connor <i>et al.</i> (1995)
Degradation by <i>P.fluorescens</i> ST subjected to inhibition by glucose, acetate and glutamate, and down-regulated by succinate and lactate.		Santos <i>et al.</i> (2000)
Metabolic integration of 3 operons for catabolism of the aromatic substrate in a functional unit (catabolon) of <i>P.putida</i> U.	phenyl acetate	Olivera <i>et al.</i> (1998)
Succinate inhibits consumption of the aromatic hydrocarbon in <i>P.putida</i> ML2.	benzene	Mason (1994)
Benzoate inhibits catabolism of phenol and acetate by <i>Ralstonia eutropha</i> . Succinate impairs benzoate consumption by blocking one key enzyme for its metabolism.	phenol, benzoate	Ampe and Lindley (1995); Ampe <i>et al.</i> (1997, 1998)
Alk pathway of the OCT plasmid of <i>P.oleovorans</i> is inhibited in rich media and rapid growth. Repressed by C-sources.	<i>n</i> -alkanes	Yuste <i>et al.</i> (1998); Staijen <i>et al.</i> (1999)
TOL pathway of plasmid pWW0 of <i>P.putida</i> mt2. Repressed by rich media, rapid growth and some carbohydrates (i.e. glucose).	toluene, <i>m</i> -xylene, <i>p</i> -xylene	Cases <i>et al.</i> (1996); Marques <i>et al.</i> (1994); Duetz <i>et al.</i> (1994, 1996, 1997); Du <i>et al.</i> (1996)
Activity of the <i>PnahG</i> promoter of <i>P.fluorescens</i> HK44 is down-regulated by rich media, glucose and toluene.	naphthalene, salicylate	Heitzer <i>et al.</i> (1994)
Catabolism of the aromatic substrate by <i>P.fluorescens</i> CA4 is inhibited by glutamate, but not by glucose or citrate.	ethyl benzene	Corkery <i>et al.</i> (1994)
<i>clc</i> pathway of <i>P.putida</i> AC27 down regulated by fumarate through inhibition of the regulator ClcR.	3Cl-benzoate	McFall <i>et al.</i> (1997, 1998)
Downregulation of the metabolism of the xenobiotic compound by glutamate, glucose and cellobiose in a <i>Flavobacterium</i> strain.	pentachlorophenol	Topp <i>et al.</i> (1988); Topp and Hanson (1990)
DMP pathway of plasmid pV1150 of <i>Pseudomonas</i> sp. CF600. Inhibited by rich media and C substrates allowing fast growth.	phenol, methyl-phenols	Sze <i>et al.</i> (1996)

promoter (Cases and de Lorenzo, 1998). Such signals include nutrient availability, but also osmolarity, temperature, chaotropic agents, contact with surfaces, and interactions with other microorganisms. Under such tough conditions, it is of essence that expression of biodegradative operons becomes tightly coupled to the physiological and metabolic state of the cells. Figure 1 shows the various types of response found in promoters that drive expression of catabolic promoters. We would predict that biodegradative operons evolve from constitutive expression to substrate-responsive and metabolically controlled transcription. In this respect, pathways found in bacteria able to degrade man-made xenobiotic compounds, either totally or partially, frequently display a range of suboptimal, non-regulated expression profiles. In contrast, biodegradation of compounds that, despite being recalcitrant, have been available to bacteria for a long time is controlled through promoters endowed with sophisticated facets to ensure the processing of substrate-specific and general physiological signals. Typical promoters of this type are those that drive biodegradation of BTEX components of petroleum (benzene, toluene, ethylbenzene, xylenes), styrene, *n*-alkanes, or side-products of wood decay such as phenylacetate, phenols and benzoates (Díaz and Prieto, 2000). Some chloro-aromatic compounds (Cl-benzoates, Cl-catechols) may also be of natural origin, thus the cognate catabolic pathways frequently show responsiveness to both the substrate and the metabolic state.

Table II shows a number of examples where the presence of a physiological control of biodegradative pathways has been observed as something superimposed on the substrate-dependent expression of the catabolic genes. Some of these will be examined in detail below. One typical environmental factor is the presence in the

same niche of alternative carbon sources, the preferential consumption of which must be decided. This may or may not be related to the overall growth rate and growth phase, a major origin of signals that can be exploited for adjustment of promoter output. The most frequent induction pattern of highly evolved catabolic promoters is that referred to as type 4 in Figure 1, which exhibits a phenomenon termed 'post-exponential induction' or 'exponential silencing'. Regardless of the mechanisms involved (discussed below), such conduct consists of a lack of transcriptional activity while bacteria grow rapidly on a nutrient-rich media, irrespective of the presence of the effector. This is then followed by rapid induction of the promoter once the growth rate of the bacteria decreases, they enter stationary phase, or cease to grow altogether. Effector-triggered post-exponential induction requires both the presence of a given effector and that cells reach a specific physiological stage. Such a stage might be defined not only by the depletion of a limiting nutrient or growth factor, but also by a particular overall energy state. To detect such a metabolic and physiological condition, and to connect it to the transcription of specific catabolic promoters, bacteria have evolved a diverse array of molecular artifices.

Assets for physiological control of catabolic operons

Unlike their eukaryotic counterparts, which seem to have a large number of transcription factors and controllable steps available to them, prokaryotic promoters have a very limited number of potential targets to integrate transcriptional co-regulation elements, namely specific regulators, sigma factors and promoter DNA. Such a paucity of molecular instruments has, however, been maximally

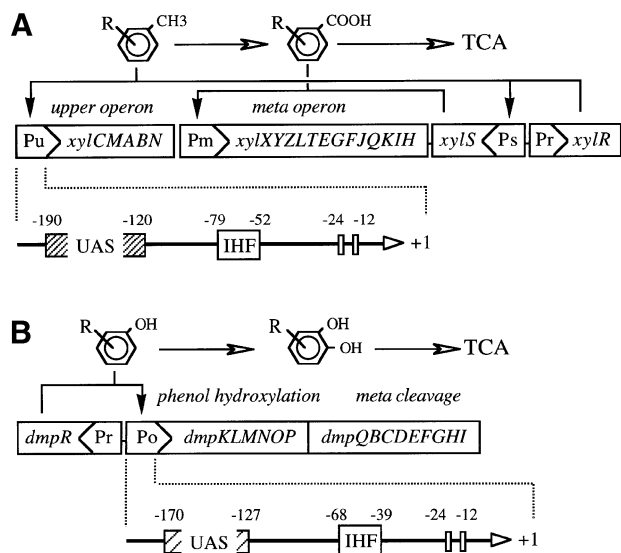


Fig. 2. Organization of the TOL and *dmp* biodegradation pathways and their cognate *Pu* and *Po* promoters. (A) The regulatory cascade of the *xyl* genes in the TOL plasmid pWW0 of *P. putida* mt-2. In the presence of *upper* pathway substrates like *m*-xylene, the *upper*-operon promoter *Pu* and the *xylS* promoter *Ps* are activated by XylR in combination with the σ^{54} -containing RNA polymerase (σ^{54} -RNAP). Subsequently, an excess of XylS product or XylS bound to its effectors (i.e. substrates of the *meta* pathway) activate *Pm*. There is no physical continuity between the *upper* and the *meta* operons. Below the scheme of the pathway, the *Pu* promoter region is expanded, showing the boundaries of relevant DNA sequences: upstream binding sites (UAS) for XylR, the -12/-24 sequences recognized by σ^{54} -RNAP, and a single IHF binding site located within the intervening region. (B) Regulation of the pV1150-encoded *dmp*-operon of *Pseudomonas* sp. CF600. The *dmpR* gene product that is responsive to phenol and cresols activates transcription of the divergently transcribed *dmp*-operon from the *Po* promoter. A subset of the *dmp* genes are involved in phenol hydroxylation, while the rest encode enzymatic activities of the *meta*-cleavage pathway for dissimilation of the catechol intermediate. The *Po* promoter region is expanded below the scheme of the *dmp* pathway. Relevant portions of the sequence are pinpointed.

combined and exploited with remarkable success. Of the many examples known (Table II) in which biodegradative pathways are subject to physiological regulation, the mechanisms involved have been studied to a significant degree in only a few instances. The diversity of regulatory strategies resulting in the same eventual phenotype is demonstrated in the cases that follow.

Metabolites that inhibit regulators

A remarkable example of how the transcriptional regulators of catabolic promoters can be subdued to the overall carbon metabolic and energetic status of cells is provided by the control of 3Cl-benzoate degradation in some strains of *P. putida*. The ortho-cleavage pathways of catechol and 3Cl-catechol are central catabolic pathways of *P. putida* that convert aromatic and chloroaromatic compounds (such as benzoate and 3Cl-benzoate) to tricarboxylic acid (TCA) cycle intermediates (McFall *et al.*, 1997, 1998). They are encoded by the evolutionarily related *catBCA* and *clcABD* operons, respectively. Expression of the *cat* and *clc* operons requires the LysR-type transcriptional activators CatR and ClcR, and the inducer molecules *cis,cis*-muconate and 2Cl-*cis,cis*-muconate.

Although the core transcriptional activation mechanisms of CatR and ClcR have been conserved in response to the presence of inducer, nature has provided some flexibility to respond to physiological signals. Transcriptional fusion studies demonstrated that the expression from the *clc* promoter is repressed when the cells are grown on succinate, citrate or fumarate, and that this repression is ClcR-dependent and occurs at the transcriptional level. The presence of these organic acids did not affect the expression from the *cat* promoter. *In vitro* transcription assays demonstrate that the TCA cycle intermediate, fumarate, directly and specifically inhibits the formation of the *clcA* transcript. No such inhibition was observed when CatR was used as an activator on either the *cat* or the *clc* template. Since both the catechol and the Cl-catechol pathways feed into the TCA cycle, but only the Cl-catechol pathway is inhibited by fumarate, there is a subtle difference in the regulation of these two pathways, where intracellular sensing of a TCA cycle intermediate leads to a reduction of chloroaromatic degradation. Titration studies of fumarate and 2-chloromuconate *in vitro* transcription assays show that the fumarate effect is concentration-dependent and reversible, indicating that fumarate and 2-chloromuconate most probably compete for the same binding site on ClcR (McFall *et al.*, 1997, 1998). This is an interesting example of the transcriptional regulation of a biodegradative pathway through the sensing of the levels of one key metabolite of the TCA cycle. Unsophisticated as it may appear, this type of metabolic downregulation of a xenobiotic-degrading pathway (*clc*) caused by a side-metabolite from a substrate that is easier to consume is probably very frequent, since just a few mutations in the targeted protein makes it amenable to a degree of physiological control.

Parasitizing sigma factors

The general transcription machinery can also be used by degradation pathways to couple expression of biodegradative operons with different physiological signals. One remarkable example is the TOL plasmid of *P. putida* mt-2 for degradation of toluene, in which the interplay of two promoters, two regulators and four sigma factors provide a very efficient control mechanism. *Pseudomonas putida* cells harbouring the TOL plasmid pWW0 are able to grow on toluene and *m-lp*-xylene as the only carbon source, owing to expression of a two-step pathway for the complete mineralization of these hydrocarbons (Ramos *et al.*, 1997). The first step (Figure 2A) involves the biotransformation of toluene/xylenes to their corresponding carboxylic acids through oxidation of one methyl group of the aromatic substrate. The second stage channels the benzoate (or toluate) into the Krebs' cycle. This follows a complex pathway summarized in Figure 2A. The biochemical steps are reflected in two separate transcriptional units, the so-called *upper* operon (encoding enzymes for oxidation of the methyl group of toluene) and the lower (or *meta*) operon (responsible for the aromatic ring fission leading to pyruvate and acetaldehyde). Expression of the *xyl* genes is tightly regulated through a complex cascade of transcriptional controls (Ramos *et al.*, 1997) that involve two regulators, the XylR and XylS proteins. These are responsible for the activation

of the *upper* and *meta* operons, respectively, thus ensuring optimal expression of the degradative activities only in the presence of pathway substrates (Figure 2A).

The *Pm* promoter (also called OP2), which drives expression of the lower operon for metabolism of benzoate and toluates all the way to the TCA cycle intermediates, is expressed at a high level throughout the growth curve (Marqués *et al.*, 1995). This transcription is dependent on the positive activator XylS (of the AraC family of prokaryotic regulators) activated by 3-methyl benzoate. Although from just inspecting the DNA sequence this promoter would be predicted to be dependent on the housekeeping sigma factor σ^{70} , recent observations (Marqués *et al.*, 1995, 1999; Miura *et al.*, 1998) have revealed a most intricate sigma succession mechanism that ensures continuous *Pm* activity throughout all stages of growth. First, it appears that thermosensitive *rpoD* mutants that transiently lack σ^{70} are still able to support *Pm* activity at the non-permissive temperature, thereby suggesting that other sigmas may actually drive promoter activity. In fact, it could be shown that it is the heat shock factor σ^H , rather than σ^{70} , that is required for *Pm* output following induction with 3-methyl benzoate. The surprising finding is that σ^H levels are generally very low unless a signal triggering the heat shock response occurs. When cells are challenged with the aromatic effector, however, it does trigger such a response, probably due to its effects on membrane properties. This is true mostly for cells that are exposed to 3-methyl benzoate (Marqués *et al.*, 1999). When cells enter stationary phase, the starvation sigma σ^S seems to take over and replace σ^H as the factor that directs *Pm* activity. It thus seems that activation of *Pm* transcription is achieved through a switch between two stress-responsive factors: σ^H in exponential phase and σ^S in stationary phase. In both cases, *Pm* is dependent on the same activator, XylS, and starts transcription in the same point. The *Pm/XylS* system reveals a strategy of coupling transcription of a specific promoter to the cell physiology by 'choosing' general stress signals mediated by sigma factors. Since the whole TOL system is plasmid encoded, it is remarkable how expression of the biodegradation functions is the result of an interplay, if not a parasitism, of general host factors with system-specific, plasmid-encoded regulators.

The σ^{54} promoter *Pu* of the TOL plasmid

In 1986, a now classic paper (Dixon, 1986) reported that the *Pu* promoter of *upper* operon of the same TOL plasmid pWW0 of *P.putida* mt-2 mentioned above (Figure 2A) had features that made its expression dependent on the *ntrA* gene. This gene was later identified as the determinant of the sigma factor for nitrogen metabolism and thus called σ^N (or, more frequently, σ^{54} ; Merrick, 1993). Since then, the σ^{54} -dependent promoter *Pu* has become a landmark for studies on the regulation of biodegradative pathways. Such extensive work has yielded detailed knowledge on both specific effector-mediated regulation and the devices that couple its performance *in vivo* to cell physiology. *Pu* is regulated by the XylR protein, which belongs to the family of prokaryotic enhancer-binding activators that act in concert with σ^{54} (Morett and Segovia, 1993; Shingler, 1996). In the presence of toluene, xylenes and other

structural analogues, the XylR protein activates the *Pu* promoter of the *upper* TOL operon (Figure 2A), using a mechanism that is generally shared by other activators of the family. This involves the binding of the regulator to upstream activating sequences (UAS) and the looping-out of the complex into close proximity with the σ^{54} -containing form or RNA polymerase bound to the $-12/-24$ region of the promoter (Morett and Segovia, 1993; North *et al.*, 1993). This event is assisted by the presence of an integration host factor (IHF)-binding site at the intervening region between the UAS and the σ^{54} -RNAP attachment site (Pérez-Martín and de Lorenzo, 1996a,b). Such an elaborate promoter architecture (Figure 2A) seems to be particularly well suited to integrating a repertoire of environmental signals.

From the early studies on the regulation of this promoter, it became evident that expression of the *upper* TOL operon was inhibited when cells grew exponentially in rich medium (Hugouvieux-Cotte-Pattat *et al.*, 1990; de Lorenzo *et al.*, 1993). This effect seemed not to require the activity of the whole complement of TOL genes, since it could be faithfully reproduced with only the regulatory elements that control transcriptional activity of *Pu*. *Pseudomonas putida* cells devoid of the TOL plasmid carrying a chromosomal insertion of the *xylR* gene and a *Pu-lacZ* fusion were unable to accumulate β -galactosidase when growing exponentially in Luria-Bertani (LB) medium, regardless of the presence or absence of *m*-xylene (de Lorenzo *et al.*, 1993). However, as soon as the cells leave the exponential growth phase and enter the stationary phase, the same *Pu-lacZ* fusion becomes extremely responsive to the aromatic inducer (de Lorenzo *et al.*, 1993; Cases *et al.*, 1996). The data of the reporter fusion match faithfully the quantitative S1 protection assays with mRNA from induced cells, so the effect certainly occurs at the transcriptional level (Marqués *et al.*, 1994). Since *Pu* is functional *in vitro* simply by mixing purified and pre-activated XylR with σ^{54} -containing RNAP and IHF (Pérez-Martín and de Lorenzo, 1996b), it is clear that additional elements, when induced, adjust transcription to the physiological state that governs the cells. These initial observations triggered a large number of studies on the mechanisms involved in such a physiological inhibition of *Pu* activity.

Various reports from different perspectives have documented that *Pu* activity is downregulated in response to exponential growth in rich media, a phenomenon referred to as catabolite repression (Duetz *et al.*, 1994, 1996, 1997; Holtel *et al.*, 1994; Marqués *et al.*, 1994), stationary-phase dependency (Hugouvieux-Cotte-Pattat *et al.*, 1990) or, as we prefer to call it, exponential silencing (Cases *et al.*, 1996) (Figure 1). At least in part, this effect can be traced to modulation of the activity of the sigma factor itself, because its overproduction shifts *Pu* derepression to an earlier growth stage (Cases *et al.*, 1996). In addition, *ftsH* mutants of *E.coli* (encoding a membrane-bound protease involved in the turnover of σ^{32}) fail to activate *Pu* (Carmona and de Lorenzo, 1999). Overproduction of σ^{54} defeats the inhibition caused by the loss of FtsH protein. Increased levels of FtsH not only restore promoter activity but also relieve the physiological inhibition of the promoter. The mechanism by which all this occurs has not yet been resolved, but may involve the action of an

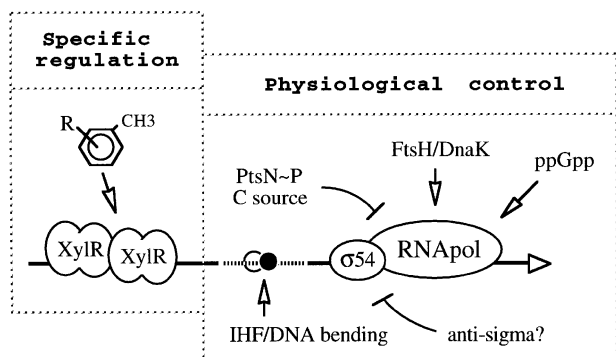


Fig. 3. Integration of specific and physiological signals on the outcome of the *Pu* promoter of the TOL plasmid pWW0. This archetypal promoter receives both inducer-specific and overall physiological inputs. The specific regulation that makes *Pu* respond to toluene and xylenes involves only the XylR regulator. On the contrary, metabolic inputs are channeled towards the transcription machinery through multiple molecular assets. These include the control of the activity or the turnover of the σ^{54} factor *in vivo* (perhaps through the FtsH/DnaK system, or an antisigma factor or both) (Cases *et al.*, 1996; Carmona and de Lorenzo, 1999), the action of the phosphorylated form of PtsN (Cases *et al.*, 1999) and also the influence of intrinsic or factor (IHF)-mediated DNA bending. The related *Po* promoter of catabolic plasmid pV1150 (see Figure 2B) seems to react positively to the intracellular levels of the alarmone ppGpp (Sze *et al.*, 1999), whereas the response of *Pu* to the stringent response is far less pronounced (Carmona *et al.*, 2000).

anti-sigma or a connection between the activity of σ^{54} and the turnover machinery of the heat shock factor σ^{32} , which involves the DnaK chaperone and the FtsH protease (Muffler *et al.*, 1997). The presence or absence of the stationary phase sigma, σ^S , does not appear to have a role in the growth dependency of *Pu* outcome. This is striking, since other metabolically regulated promoters (see below) benefit from this global starvation and growth phase signal for behaving in a fashion indistinguishable from that observed in *Pu*. Finally, it is remarkable that in spite of being strongly inhibited by casamino acids, the *Pu* promoter functions normally in a *relA/spoT* strain of *E.coli* that fails to accumulate the alarmone (p)ppGpp that is typical of the stringent response (Cashel *et al.*, 1996; Carmona *et al.*, 2000). Yet, overproduction of RelA (predicted to force an increase ppGpp levels) does increase *Pu* activity and causes a certain relief of its exponential silencing (Carmona *et al.*, 2000).

Carbon sources versus metabolic state

Besides the effect of growth rate and growth phase, the presence of certain carbon compounds such as glucose or gluconate also restrain *Pu* activity (Holtel *et al.*, 1994; Cases *et al.*, 1999). It should be noted that although a regulator analogue of the well characterized catabolite repression protein (CRP) of *E.coli* does exist in pseudomonads (named Vfr), it appears to play no role in catabolite repression, but instead is involved in quorum sensing control (Collier *et al.*, 1996; Albus *et al.*, 1997). Even transport of glucose (a nearly universal inducer of catabolite repression) is completely different in *Pseudomonas* (Collier *et al.*, 1996). Catabolism of this sugar requires its predominant extracellular transformation into gluconic acid, followed by the induction of a specific

energy-dependent gluconate uptake system. Once internalized, gluconate is phosphorylated and entered into the central metabolism as 6-phosphogluconate (Schleissner *et al.*, 1997). These peculiarities in the carbon metabolism of *P.putida* make it unlikely that the inhibition of *Pu* activity by glucose and gluconate takes place through a typical cAMP/CRP-mediated mechanism of the sort so well known in *E.coli* (Kolb *et al.*, 1993). Furthermore, the C-source dependent inhibition of *Pu* activity occurs within a different range than that caused by rapid growth. While exponential silencing in rich medium completely abolishes *Pu* activity, glucose does not inhibit promoter output by more than two-thirds of the maximal activity. The gene *ptsN*, encoding a new member of the phosphoenolpyruvate:sugar phosphotransferase system (PTS) (Postma *et al.*, 1993; Saier, 1996; Saier *et al.*, 1996), has been related to this modulation, since its loss alleviates glucose inhibition of *Pu* in *P.putida* (Cases *et al.*, 1999). Glucose assimilation is not affected in this mutant, suggesting that this gene participates in sensing rather than metabolizing the carbohydrate. Interestingly, *ptsN* mutants are still subject to exponential silencing. In the other direction, cells relieved of such silencing because of σ^{54} overproduction remain repressible by glucose (Cases and de Lorenzo, 2000). These observations indicate that the growth-rate inhibition of *Pu* and the downregulation by glucose can be separated genetically and, surely, are channeled into the promoter via different pathways (Figure 3). The mechanism by which PtsN exerts its inhibition on *Pu* in the presence of glucose is not yet known, but may involve other proteins of the PTS family such as PtsO (Powell *et al.*, 1995; I.Cases and V.de Lorenzo, unpublished). The whole picture is by no means complete and will remain the subject of further investigation.

Variations on the σ^{54} promoter theme

Even promoters that are organized with the same control elements and that show similar physiological regulation patterns have evolved a different regulatory mechanism, as the *Po/DmpR* system illustrates. *Po*, the phenol-responsive promoter that drives expression of the single *dmp* (dimethylphenol) operon of plasmid pV1150 harboured by *Pseudomonas* sp. CF600 (Powlowski and Shingler, 1994) (Figure 2B) is, as *Pu*, dependent on σ^{54} and is activated by a cognate protein named DmpR. Although responsive to phenol and methyl-phenols rather than xylenes, the sequence of this protein is largely similar to that of XylR (Shingler *et al.*, 1993). In fact, it was soon proven that the major domains of the two proteins could be swapped without a loss of function and that either protein could activate the other cognate promoter and recognize their mutual UAS (Fernández *et al.*, 1994; Shingler and Moore, 1994). This suggested that XylR and DmpR behaved more as variants of the same protein than as different proteins. When *in vivo* physiological studies of the *Po* promoter were carried out, it also became clear that *Po* was subjected to exponential silencing, i.e. no transcription could be detected while cells grew rapidly in rich LB medium (Sze *et al.*, 1996). Although such behaviour would superficially be identical to that reported for *Pu*, further insights have revealed that the mechanisms involved differ significantly between the XylR/*Pu* and the

DmpR/*Po* systems. For instance, Sze *et al.* (1996) reported that the exponential silencing observed in *Po* was, in general, inversely proportional to growth rate. Thriving in certain carbon sources thus inhibits promoter activity to the same extent that it increased growth rate. In a subsequent step, it was shown that cells lacking *relA* and *spoT* (Cashel *et al.*, 1996; see above) failed to activate the σ^{54} promoter *Po* at any growth phase (Sze and Shingler, 1999). On the contrary, overexpression of *relA* (and the ensuing increase in ppGpp levels) entirely relieved exponential silencing. These observations suggested that *Po* activity was dependent on the intracellular accumulation of ppGpp related to growth rate, and provided a rationale to understand the coupling of this σ^{54} promoter to metabolic signals related to starvation. The notion has been strengthened by the observation that some RNAP variants, which bear mutations that mimic the response to ppGpp, overcome exponential silencing as well (Sze and Shingler, 1999). Such a subordination to ppGpp is in contrast to the lesser dependence of the XylR/*Pu* system to the same signal (Carmona *et al.*, 2000; see above), and this difference is informative. *In vitro* transcription of the *Po* promoter with purified XylR was highly dependent on ppGpp, whereas *Pu* in the same experiment could be transcribed without any addition of the nucleotide (Carmona *et al.*, 2000). On the basis of comparing the mechanisms that produce the physiological silencing of *Pu/XylR* with those of *Po/DmpR*, it appears that the class of metabolic signals that end up interweaving the control of such σ^{54} promoters is determined in part by their very DNA sequence.

Optimizing DNA sequence as a regulatory framework

A final facet of the physiological control of some σ^{54} systems is the role of the very promoter DNA sequence as a possible port of entry of additional signals from cell metabolism. Since the early observation that *Pu* bore a functional binding site for IHF (Figure 2A) that is essential to maintain a suitable promoter architecture, DNA bending was considered a potential regulatory obstacle for the system (Pérez-Martín and de Lorenzo, 1995, 1997a). It is known that IHF accumulation in *E.coli* is growth-phase-dependent (Aviv *et al.*, 1994), so the bend produced by the factor could contribute to the enhanced activity of *Pu* in stationary phase. However, this notion was discarded because *Pu* promoter variants in which the IHF binding site had been replaced by a statically bent DNA sequence, and was thus IHF independent, were still subject to exponential silencing (Cases *et al.*, 1996). In addition, it appeared that in *E.coli*, the nucleoid-associated protein HU can replace the DNA-bending function of IHF (Pérez-Martín and de Lorenzo, 1997b), so that the effect of the loss is more difficult to monitor *in vivo*. But following these observations in *E.coli*, it was noticed that a *P.putida* IHF knockout mutant failed entirely to support any *Pu* activity (Calb *et al.*, 1996). This was surprising, since there is a functional analogue of HU in *P.putida* (F.Bartels and V.de Lorenzo, unpublished), and thus some promoter function should be expected. A clue to understanding this phenomenon was the realization that IHF was necessary to recruit the σ^{54} -RNAP holoenzyme to the promoter DNA

through a mechanism that involved the interactions of the C-terminal domain of the α subunit of the RNAP (α CTD) with an upstream DNA, UP-like sequence (Bertoni *et al.*, 1998). Under limiting concentrations of σ^{54} -RNAP, such a recruitment becomes a kinetic barrier to promoter activity, more limiting than XylR activation itself (Carmona *et al.*, 1999). On the basis of these observations, the role of IHF in physiological control will have to be re-assessed. Whether such a possible regulatory device is also relevant in other σ^{54} -dependent promoters that contain IHF sites (Dworkin *et al.*, 1998; Wassem *et al.*, 2000) remains unknown.

All the data collected so far on the physiological control of σ^{54} promoters suggests that they are particularly well suited to group a whole range of metabolic symptoms into one or more steps of the transcription initiation process. With only two constant players (the substrate-responsive UAS-binding regulator and σ^{54}), these promoters afford a stunning amalgam of the effector-specific transcription with the cell physiology. The specific choice of physiological signals (coupling to the stringent response, to the turnover of other sigma factors, to the PTS system of C-sensing, to the physiological levels of IHF, to growth phase-dependent DNA bending, etc.) seems to be mostly determined by the diversity of DNA sequences at the promoter. Not unlike a music score instructing the entrance of specific instruments to play a melody, in the promoters discussed above, it becomes manifest that distinct instruments can end up performing the same theme.

Promoters responding to gasoline?

While the *Pu/XylR* and *Po/DmpR* systems have developed virtually the same behaviour by working upon variations of the σ^{54} theme, a sophisticated degree of physiological control can be generated as well with quite different molecular assets. The OCT (octane degradation) plasmid borne by *Pseudomonas oleovorans* Gp01 allows this microorganism to grow on medium chain-length *n*-alkanes (van Beilen *et al.*, 1994). This is due to a plasmid-borne metabolic pathway, whose induction is regulated by the AlkS protein (of the LuxR/MaIT family of transcriptional regulators) and the expression of which is fully σ^{54} independent. In the presence of alkanes or the gratuitous inducer dicyclopentyl ketone (DCPK; see Table I), AlkS activates the AlkB promoter, which drives expression of most genes of the pathway. The study of the behaviour of chromosomally inserted *PalkB-lacZ* fusions and S1 nuclease protection assays revealed that the promoter remains silent when cells grow exponentially in rich LB medium, even if cultures are exposed to the inducer (Yuste *et al.*, 1998). A milder downregulation occurred also when cells were grown in various carbon sources such as lactate or succinate (Yuste *et al.*, 1998; Staijen *et al.*, 1999). Phenomenologically, these inhibitory effects linked to exponential growth and carbon sources were not unlike those seen in the *Pu* and *Po* promoters above. However, given the very different promoter structures, it could be anticipated that the mechanisms involved ought to be different. An intriguing similarity with DmpR and PhlR, though (see above), is that overproduction of AlkS seemed

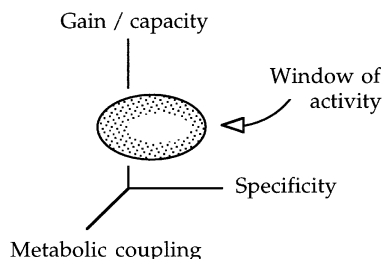


Fig. 4. Evolutionary optimization of promoter performance. The scheme illustrates the black cat/white cat principle discussed in the text. Promoters recruited to respond to novel environmental signals, such as unfamiliar carbon sources, gravitate necessarily towards an optimal window of activity. Such a window is defined as the combination of: (i) the various working ranges of promoter gain/capacity (inducibility versus absolute promoter output); (ii) the specificity of the regulator/promoter pair for a given chemical; and (iii) the connection of promoter activity to the cell physiology. The drift towards such an optimal window is independent of the molecular mechanism to reach it. The presence of specific DNA/promoter/regulators combinations in front of given biodegradative operons thus reflect much more the evolutionary history of the system than a requirement for a distinct type of transcriptional factor.

to overcome the silencing imposed by rapid growth in rich medium (Müller *et al.*, 1996; Yuste *et al.*, 1998).

An interesting edge to the control of *PalkB* expression came through the realization that *alkS* transcription was itself regulated by growth phase. This is because the *PalkS* promoter seems to have a preference for the RNAP holoenzyme bearing the starvation sigma σ^S (Canosa *et al.*, 1999). Although this could be related to the physiological control of *alk* genes, it might deal instead with a separate aspect of the system. More recent studies by Rojo's group have shown that the *PalkS* promoter is more complex (Canosa *et al.*, 2000). The promoter region includes *PalkS1*, which is fully σ^S -dependent and thus silent during exponential growth and active in the stationary phase. In the absence of alkanes, the AlkS protein represses *PalkS1* moderately, thus achieving a reasonable level of expression of the regulator. But when cells face the inducer, AlkS both completely represses *PalkS1* and activates a second, downstream promoter (*PalkS2*). In this manner, activation of *PalkS2* allowed a very efficient transcription of *alkS* when alkanes are in the medium. On this basis, it is plausible that AlkS levels are controlled by a positive feedback mechanism, which leads to a rapid increase in *alkS* transcription when alkanes are present and causes a fast off-switch when alkanes are depleted. Yet, it happens that *PalkS2* itself is inhibited by exponential growth in rich medium and is downregulated by some carbon sources such as lactate, a phenomenon that still deserves further investigation (Canosa *et al.*, 2000).

Black cat/white cat: evolutionary convergence of physiological regulation

In spite of the very different regulatory schemes present in the *alk* system of the OCT plasmid as compared with the σ^{54} -dependent routes of pWW0 and pV1150, it is amazing how both have evolved via entirely disparate mechanisms to achieve the same result: (i) responding to specific effectors; (ii) reacting to the presence of other carbon

sources in the medium; and (iii) weaving promoter activity to the growth phase. The first is achieved through different regulators (AlkS and XylR/DmpR) belonging to completely unrelated families of proteins, the second through a connection of an undisclosed catabolite repression mechanism to proteins of the PTS system, and the third by recruiting stationary phase or stress sigmas, by coupling polymerase activity to ppGpp or by checking the activity of the sigmas by additional growth-phase-dependent factors. But these considerations, as discussed above, could also be extended to each of the other catabolic promoters where the issue has been studied and that involve other strategies for coupling promoter activity to the cell physiology.

On the basis of the comparisons between the regulatory systems presented in this review, we entertain the notion that catabolic pathways for recalcitrant compounds (or, by extension, for any new compound) evolve in three major planes that define its window of activity in the environment (Figure 4). First, it is the assembly of the complement of catabolic enzymes that are required to obtain a suitable energy return out of the metabolism of a given chemical. Since the number of biochemical strategies to break bonds is limited, this step will generally follow quite strict rules imposed by the very chemistry of the biodegradation process. Secondly, newly developed metabolic operons mature from constitutive or semi-constitutive expression to conditional transcription responsive to the presence of the substrate in the medium (Figure 1). The unfolding of substrate specificity may start by enrolling the residual responsiveness of an already existing promoter-regulator pair, so the early control stages are necessarily relaxed. The choice of a given promoter-regulator pair may depend much more on the evolutionary story of the degradation operon(s) than on the specific properties of the regulatory proteins; it is indeed frequent to find very similar biodegradative operons controlled by very different promoter types (van der Meer *et al.*, 1992). But thirdly, in the highly competitive environment that predominates in sites polluted with recalcitrant compounds, promoters must be able to process a range of simultaneous environmental signals so that expression of a degradation pathway for one compound present in the medium does not turn against the overall ecological performance of the host. To this end, it is of essence that the transcription machinery of specific promoters are able to sense the physiological state of the cells as a whole and react to it. That refining specific regulation and coupling it to cell physiology are frequently overlapping processes in time can be noted by comparing Tables I and II. Whether sensing of that kind occurs through responding to a metabolite of the TCA cycle, to a nutritional alarmone, to a stress signal or to proteins responding to extra carbon sources does not really seem to matter, provided that it links one thing to the other.

This molecular black cat/white cat principle applied to biodegradative pathways has repercussions that extend beyond the mere basic understanding of bacterial promoters. It has major consequences in the way novel expression systems might be designed for biotechnological processes (Schroetckh *et al.*, 1999; Farmer and Liao, 2000), particularly in those applications involving metabolic engineering projects of bacteria destined for the environment as bioremediation agents (de Lorenzo, 1994;

Timmis *et al.*, 1994; Matin *et al.*, 1995; Timmis and Pieper, 1999).

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