

Convergent evolution of similar enzymatic function on different protein folds: The hexokinase, ribokinase, and galactokinase families of sugar kinases



PEER BORK, CHRIS SANDER, AND ALFONSO VALENCIA

European Molecular Biology Laboratory, Heidelberg, Germany

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Abstract

Kinases that catalyze phosphorylation of sugars, called here sugar kinases, can be divided into at least three distinct nonhomologous families. The first is the hexokinase family, which contains many prokaryotic and eukaryotic sugar kinases with diverse specificities, including a new member, rhamnokinase from *Salmonella typhimurium*. The three-dimensional structure of hexokinase is known and can be used to build models of functionally important regions of other kinases in this family. The second is the ribokinase family, of unknown three-dimensional structure, and comprises pro- and eukaryotic ribokinases, bacterial fructokinases, the minor 6-phosphofructokinase 2 from *Escherichia coli*, 6-phosphotagatokinase, 1-phosphofructokinase, and, possibly, inosine-guanosine kinase. The third family, also of unknown three-dimensional structure, contains several bacterial and yeast galactokinases and eukaryotic mevalonate and phosphomevalonate kinases and may have a substrate binding region in common with homoserine kinases.

Each of the three families of sugar kinases appears to have a distinct three-dimensional fold, since conserved sequence patterns are strikingly different for the three families. Yet each catalyzes chemically equivalent reactions on similar or identical substrates. The enzymatic function of sugar phosphorylation appears to have evolved independently on the three distinct structural frameworks, by convergent evolution. In addition, evolutionary trees reveal that (1) fructokinase specificity has evolved independently in both the hexokinase and ribokinase families and (2) glucose specificity has evolved independently in different branches of the hexokinase family. These are examples of independent Darwinian adaptation of a structure to the same substrate at different evolutionary times. The flexible combination of active sites and three-dimensional folds observed in nature can be exploited by protein engineers in designing and optimizing enzymatic function.

Keywords: enzymatic function; galactokinase; hexokinase; ribokinase; substrate specificity evolution; sugar kinases

Sugars are used by cells as a source of energy or carbon. The first step in sugar metabolism after transport into the cell is phosphorylation catalyzed by specific sugar kinases. All metabolic sugars are phosphorylated to trap them inside the cell and to prepare them for further chemical reactions, either catabolic or anabolic. The main catabolic pathway is glycolysis, with glucose as the key sugar. The other key sugar is ribose, essential in nucleotide biosynthesis. Other abundant sugars are fructose and galactose, whereas ribulose, xylulose, or fucose and others are rare and can only be processed by specialized microorganisms.

Advances in sequencing technology have led to a rapidly increasing number of available primary structures.

In combination with recent biochemical data, this sequence information is used here to obtain some insight into the evolution of this important class of enzymes and into functionally and structurally important sequence regions. A detailed understanding of sugar kinases is particularly important in the context of human hereditary diseases, glycerol kinase deficiency (Seltzer et al., 1989), and non-insulin-dependent diabetes mellitus (Hattersley et al., 1992; Stoffel et al., 1992; Vionnet et al., 1992).

Results

The hexokinase family

Hexokinase has an ATPase domain with the same basic fold and active site as that of actin and the hsp70 family

Reprint requests to: Protein Design Group, European Molecular Biology Laboratory, Meyerhofstrasse 1, D-6900 Heidelberg, Germany.

of heat shock proteins (Bennet & Steitz, 1980; Flaherty et al., 1990, 1991; Kabsch et al., 1990; Holmes et al., 1993). Probable additional members of the family can be identified by sequence pattern searches. These include various sugar kinases as well as the prokaryotic cell cycle proteins Mreb, Ftsa, and Stba (Bork et al., 1992). The ligand ATP is located between two main domains, and its phosphate tail is bound by two beta-hairpins, one from either domain (hexokinase differs slightly from actin and hsc70 in

one of the phosphate-binding beta-hairpin loops). The adenosine base is bound by a nearby loop, and two contacting helices appear to form a hinge region between the two main domains (see kinemages).

Diverse specificities

The hexokinase family (Table 1) includes eukaryotic hexokinases and glucokinases as well as prokaryotic gluco-, ribulo-, glucono-, xylulo-, glycerol-, fructo-, rhamno-, and

Table 1. The sugar kinase families

Sequence ^a	E.C.	Pathway ^b	Sequence code ^c
Hexokinase-like		Mainly catabolic	
Hexo-	2.7.1.1	Glycolysis	HXKA_YEAST, HXKB_YEAST, HXKA_MOUSE, HXKB_HUMAN, HXKB_RAT, HXK2/RAT, HXK3/RAT, HXST/HUMAN, HXKH/HUMAN, HXKP/MOUSE, HXKC/BOVIN
Gluco-	2.7.1.2	Glycolysis	HXKG_ZYMMO, HXKG/STRC, HXKG/HUMAN, HXKH_RAT ^d , HXKP_RAT, HXKG_YEAST
Fructo-	2.7.1.4	Glycolysis	FRK/ZYMMO
Ribulo-	2.7.1.16	Pentose cycle	KIRI_ECOLI, KIRI_SALTY
Glucono-	2.7.1.12	Pentose cycle	GNTK_BACSU
Xylulo-	2.7.1.17	Pentose cycle	XYLK_ECOLI, XYLK_LACPE, XYLK_ACTMI, XYLK/STRU, XYLK/ARTBA, XYLK/THERM
Glycero-	2.7.1.30	Glycolysis	GLPK_BACSU, GLPK_ECOLI
Fuco-	2.7.1.52	Pentose cycle	FUCK_ECOLI
Rhamnulo-	2.7.1.5		RHAK/SALTY
Ribokinase-like		Several roles	
Fructo-	2.7.1.4	Glycolysis, sucrose	SCRK_VIBAL, SCRK_SALTY, SCRK/KLEPN
Ribo-	2.7.1.15	Pentose cycle, nucleotide metabolism	RBSK_ECOLI, RBSK_YEAST, RBSK/PACDE
6-Phosphofructo-Minor enzyme ^e	2.7.1.11	Glycolysis?	K6P2_ECOLI
6-Phosphotagato-		Glycolysis?	K6PT/STAAU (LACC_STAAU), K6PT/STRMU, K6PT/LACLA (LACC_LACLA)
1-Phosphofructo-	2.7.1.56	Glycolysis	KIPF_ECOLI, KIPF_RHOCA, KIPF_XANCP
Inosine-guanosine-	2.7.1.73	Purine metabolism	INGK_ECOLI
Galactokinase-like		Mainly anabolic	
Galacto-	2.7.1.6	Lactose cycle	GAL1_ECOLI, GAL1_KLULA, GAL1_SACCA, GAL1_SALTY, GAL1_STRLI, GAL1/LACHE
Mevalonate-	2.7.1.36	Cholesterol synthesis	KIME_RAT, KIME_YEAST
P-mevalonate-	2.7.4.2	Cholesterol synthesis	ERG8_YEAST
Homoserine- ^f	2.7.1.39	Threonine synthesis	KHSE_BACSU, KHSE_BRELA, KHSE_CORGL, KHSE_ECOLI, KHDE_FREDI, KHSE_YEAST

^a For the names of the sugar kinases, append the word kinase after the hyphen.

^b Metabolic pathways in which the enzymes are active, with a question mark where the pathway is very uncertain.

^c Sequence codes are from SWISSPROT where available and have an “_” between enzyme and species abbreviation (Bairoch & Boeckmann, 1992). Other sequences were taken from the literature (see Materials and methods for references), with names chosen by us and “/” as the separator between enzyme name and species. Detailed information for each SWISSPROT sequence including references appears on the Diskette Appendix. The codes for the species are: ACTMI, *Actinoplanes missouriensis*; ARTBA, *Artherobacter*; BACSU, *Bacillus subtilis*; BRELA, *Brevibacterium lactofermentum*; CORGL, *Corynebacterium glutamicum*; ECOLI, *Escherichia coli*; FREDI, *Fremyella diplosiphon*; KLEPN, *Klebsiella aerogenes*; KLULA, *Kluyveromyces lactis*; LACHE, *Lactobacillus helveticus*; LACLA, *Lactococcus lactis*; LACPE, *Lactobacillus pentosus*; RHOCA, *Rhodobacter capsulatus*; SACCA, *Saccharomyces carlsbergensis*; SALTY, *Salmonella typhimurium*; STAAU, *Staphylococcus aureus*; STRC, *Streptomyces coelicolor*; STRLI, *Streptomyces lividans*; STRMU, *Streptococcus mutans*; STRU, *Streptomyces rubiginosus*; THERM, *Thermus thermophilus*; VIBAL, *Vibrio alginolyticus*; XANCP, *Xanthomonas campestris*; ZYMMO, *Zymomonas mobilis*.

^d The glucokinase sequence Hxkh_Rat differs from the second isoenzyme, Hxkp_Rat, only in the first 15 amino acids due to differential splicing of the gene. Hxkp_Rat is therefore omitted from the tree (Fig. 2).

^e The minor isoenzyme 2 of 6-phosphofructokinase from *E. coli* has only about 10% in vitro activity compared to the major, structurally unrelated, counterpart (Daldal, 1984).

^f Homoserine kinase shares only one conserved region with the other enzymes of this family (Fig. 5; for interpretation see text).

fucokinases. In spite of their different substrate specificities, most of the prokaryotic sugar kinases have clear overall similarities to each other (Fig. 2, right), detectable by standard homology search programs like Fasta (Pearson & Lipman, 1988). Two members difficult to detect are ribulokinase, with only about 20% identical residues relative to other members of this branch, and rhamnokinase, which was identified here by a combination of alignment and pattern searches. The member closest to rhamnokinase is fucokinase (25% amino acid identity within 409 residues; Fasta score of 266; Figs. 1, 2).

Three-dimensional models

Based on common sequence patterns in the structural core region (Bork et al., 1992; Fig. 1), a plausible three-dimensional model can be built of the core of any protein in this family, based on the structure of yeast hexokinase. Using such a model, structural and functional information can be transferred from one member of the family to the others, even when sequence similarity is low. This can be extremely useful for the identification of residues that participate in active sites or in the interpretation of mutant phenotypes. For example, Stoffel et al. (1992) used a model of human glucokinase to suggest that the possible role two point mutants, in residues Thr-228 and Gly-261, in early-onset non-insulin-dependent diabetes mellitus is to disrupt ATP and glucose binding.

Evolutionary tree

An evolutionary tree (Fig. 2) of the hexokinase family has several interesting features. The major division is into

eukaryotic and prokaryotic enzymes. Within the prokaryotic subfamily there is a very wide spread of specificities to different sugars (Fig. 2, right and middle). The eukaryotic enzymes of this family (Fig. 2, left) are hexokinases (type I, II, and III isoenzymes) and glucokinases (type IV hexokinases). The mammalian hexokinases have about twice the molecular weight of all other sugar kinases of this family and contain two similar domains, probably as a result of gene duplication (Easterby, 1971; Nishi et al., 1988; Schwab & Wilson, 1989). Within one species, the mammalian glucokinases differ mainly at their N-terminal 15 amino acids as a result of differential splicing.

Nt and Ct domains of mammalian hexokinases – Functional or regulatory?

The eukaryotic members form four separate clusters (Fig. 2, left): the C-terminal domains of mammalian hexokinases, their N-terminal domains, mammalian glucokinases, and yeast sequences; each of these, except for the N-terminal domain of rat hexokinase type III, has all the residues identified, by analogy, to be involved in ATP binding (Bork et al., 1992). However, biochemical experiments indicate that the N-terminal domain is regulatory and the C-terminal domain catalytic (White & Wilson, 1989). Unfortunately, the separation in the tree of N-terminal and C-terminal domains of mammalian hexokinases, although indicative of a divergence of function, throws no further light on the possible regulatory role of the N-terminal domain.

HXKB_YEAST	81	DFLAIDLGGTNRVVLVVLKGG	-46-	SEPIPLGTFSPFASQNKIN	-37-	VVALINDTTGTLVASYYTD	-1-	ETKMGIIFGTGVNGAYYD		
HXKG_YEAST	80	VLLAADLGGTNRFRICSVNLHG	-53-	AKPMKLGTFSPVVDQTSLN	-38-	VVALTNDTVGTYLSHCYTS	-12-	EPVIGCIFGTGTNGCYME		
HXKH_RAT	73	DFLSLDLGGTNRVRLVVKVGG	-47-	HKKLPLGTFSPVVRHEDLD	-38-	VVAVMNDTVATMISCYEED	-1-	QCEVGMIVGTGCNACYME		
HXKA_MOUSE1	79	DFIALDLGGSSFRILRVQVNH	-45-	DKKLPVGTTFSPPCRQSKID	-38-	IVAVVNDTVGTMTCGYDD	-1-	QCEVGLIIGTGTNACYME		
HXKA_MOUSE2	527	DFIALDLGGTNRVRLVVKIRG	-45-	GPRMPLGTFSPFCKQTSLD	-38-	VVAVVNDTVGTMTCAYEE	-1-	SCEIGLIVGTGSNACYME		
HXKG_ZYMNO	2	EIVALDIGGTHARFSAIEVSS	-57-	NPWVLRPATLNEKLD	.IID	-0-	THVLINDFGAVAHAVAHMD	-17-	GVITILGPGTGLVGAHLL	
Kiri_Salty	3	IAIGLDFGSDSVRALAVDCAT	-54-	ANVVIGVDSSTGSGTP	.APID	-171-	ISGGAFDCHMGAVGAGA	.Q	-0-	PNTLVKVIPTSTCDILIA
Glpk_Ecoli	5	YIVALDQGTSSRAVMDHDA	-46-	DQIAAIGITNQRETT	.IVWE	-149-	ISGIAGDQQAALFGQLCVK	-0-	EGMARNTYGTGCFMLMNT	
Xylk_Ecoli	1	MYIGIDLGTSGVKVILLNEQG	-43-	QDVKALGIAGQMHA	.TLLD	-142-	VVAGGGDNAAGAVGVGMVD	-0-	ANQAMLISLGTSGVYFAVS	
Gntk_Bacsu	4	YMLGIDIGTSTKAVLFSENG	-43-	KRISFISFSSAMHSV	.TAID	-145-	FVIGASDGVLSNLGVNAIK	-0-	KGEIAVTIGTSGAIRTII	
Fuck_Ecoli	15	VILVLDGCGATNRAIAVNRQG	-45-	CHIRGIAVTTFPGVDG	.ALVD	-145-	VISAGHDTQFALFGAGA	.E	-0-	QNEPVLSSGTWEILMVR
Rhak/Salty	5	HCVAVDLGASSGRVMLARYDS	-45-	ILDISIGIDTWGVDY	.VLLD	-140-	VAVASHDTASAVIASPLAN	-0-	KHSAYLSSGTWLMGFES	
		phosphate 1		sugar binding?		connect 1		phosphate 2		
HXKB_YEAST	-144-	IGARAARLSVCGI	-7-	GKVTGHIAADGSVYNYRYPGFKEKAANALKDIY	-13-	IVPAEDGSGAGA AVIAALAAQ				
HXKG_YEAST	-152-	ISRRSAYLA AVPL	-13-	YHGEVEIGCGDSVVEYYPGFRSMLRHALLSP	-11-	LKIAKDGSGVGAALCALVA*				
HXKH_RAT	-138-	VSTRAAHMCSAGL	-13-	DVMRITVGVGDSVYKLPSPKRFPHASVRRLT	-5-	FIESEEGSGRGAALVSAVAC				
HXKA_MOUSE1	-138-	VSFRAANLVAATL	-13-	PRLRITVGVGDSVYKLPSPKRFPHASVRRLT	-5-	FLLSESGSGRGAAMVAVAY				
HXKA_MOUSE2	-138-	VSKRAALCGAGM	-13-	DHLNVTVGVGDTLYKLPSPKRFPHASVRRLT	-5-	FLLSEDSGSGRGAALITAVGV				
HXKG_ZYMNO	-34-	RFRRVSIERIIISG	-4-	NIYEALAAIEGVFPFSLDDIKLWQMALEGKDN	-21	LAQQRTSVVIGGGVGLRIAS				
Kiri_Salty	-131-	FGARAIQECFTDQ	-0-	GIAVNNVMALGGIARKNQVIMQVCCDVLNRPL	-1-	IVASDQCCALGAAIFA AVAA				
Glpk_Ecoli	-89-	HIIRATLESIA YQ	-12-	GIRLHALRVDDGAVA	.NNFLMQFQSDILGTRV	-1-	RPEVREVTALGAAYLAGLAV			
Xylk_Ecoli	-87-	ELARAVLEGVGYA	-11-	GIKPQSVTLIGGGAR	.SEYWRQMLADISGQQL	-2-	RTGGDVG PALGAARLAQIAA			
Gntk_Bacsu	-97-	HMIRAALLEGVIYN	-12-	DGPVTRIQATGGFAR	.SEVWRQMMSDIFESEV	-1-	VPESYESSCLGACILGLYAT			
Fuck_Ecoli	-83-	HFYRAALEGLTAQ	-12-	HFKAPELLLVGGGSR	.NTLWNQIKANMLDIPV	-1-	VLLDAAETTVAGAALFGYGV			
Rhak/Salty	-88-	ELARCFDLSLALL	-12-	GEKFTQLHIVGGGCQ	.NSLLNQLCADACGIRV	-0-	MAGPVEASTLGNIGIQMLT			
		helix		adenosine		connect 2				

Fig. 1. Hexokinase family: aligned conserved regions of a representative subset of family members. Numbers between the blocks indicate the length of intervening segments. Protein names indicate the name of the protein followed by a species identifier, taken from SWISSPROT when available (e.g., Hxkb_Yeast) or chosen by us (e.g., Rhak/Salty). A list of all the enzymes used is in Table 1. Bottom line: functional and structural identification of the conserved regions, in analogy to the known 3D structure of yeast hexokinase: phosphate 1 and 2 and adenosine are involved in ATP binding, connect 1 and connect 2 in putative hinge motion (Bork et al., 1992); sugar binding corresponds to a possible glucose-binding site in hexokinase (Arora et al., 1990); and the helix is an additional conserved region common to the hexokinase-like sugar kinases.

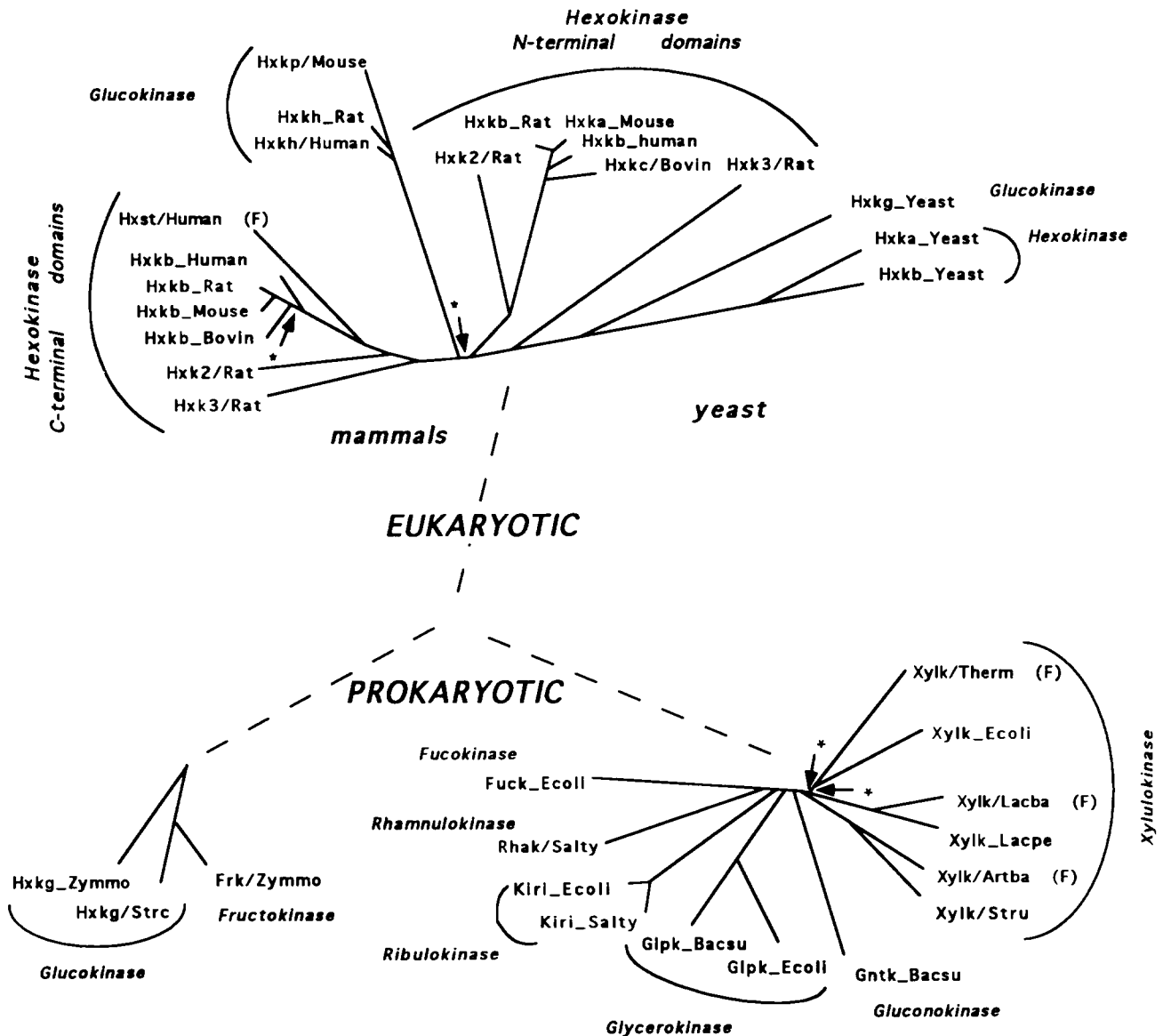


Fig. 2. Hexokinase family: evolutionary tree of all currently available family members. The distances from the root of the tree are undetermined (dotted lines) as the similarity between the three subfamilies is limited to the seven conserved boxes. The branch lengths are roughly proportional to the amount of sequence difference, as calculated using the program Clustal V (Higgins et al., 1992) with full-length sequences as input. An asterisk marks lack of confidence in the precise branching order (confidence less than 95% in the bootstrapping option of Clustal V). (F) indicates sequence fragments. The N-terminal and C-terminal domains of mammalian hexokinases, a probable result of gene duplication, are entered separately (left side of figure). All proteins in the tree probably have a fold similar to that of yeast hexokinase and acts as well as of the ATPase domains of hsp70 heat shock proteins and some prokaryotic cell cycle proteins (Bork et al., 1992). The evolutionary distances (sums of branch lengths) between the xylulokinases of different species are as large as those between the xylulokinases and enzymes of other substrate specificities (right) indicating that substrate specificity is only one of several evolutionary constraints governing sequence variation. The presence of glucokinases in three separate branches (two in the left subfamily; one in the center one) is suggestive of multiple independent evolution of this specificity. Note also that fructokinase activity (center of figure) appears to have evolved independently in this structural family—fructokinases with a different fold are in Figure 4. Notation of protein names as in Table 1.

Independent evolution of glucose specificity

An interesting feature of the tree (Fig. 2) is the fact that glucokinases appear in three clusters in separate branches: (1) mammalian glucokinases form one cluster (Fig. 2, far left), (2) yeast glucokinase appears to cluster

with yeast hexokinases rather than with mammalian glucokinases (Fig. 2, left), and (3) bacterial glucokinases from *Zymomonas mobilis* and *Streptomyces coelicolor* are grouped with *Zymomonas* fructokinase (Fig. 2, middle). A divergent evolutionary relation between these

apparently disconnected clusters is unlikely. Rather, evolutionary convergence to glucose specificity independently in mammals, yeast, and bacteria probably must be invoked. It is unlikely, however, that identical active site constellations are responsible for glucose specificity in the three cases. Convergence may be limited to similarity in the ability to bind glucose specifically and may not extend to the precise way in which this is achieved. Overall, the tree of hexokinase-related sugar kinases present a remarkable diversity of specificities (Table 1), all achieved by fine tuning of specific regions in the same basic three-dimensional framework.

The ribokinase family

The second structurally distinct family is composed of prokaryotic sequences related to ribokinase (Table 1), including several fructokinases, the minor 6-phosphofructokinase from *Escherichia coli*, 1-phosphofructokinases, and 6-phosphotagatokinases, with some similarities already described (Daldal, 1983, 1984; Blatch et al., 1990; Orchard & Kornberg, 1990; van Rooijen et al., 1991) and reviewed by Wu et al. (1991). Here we have attempted to assemble a complete set of the currently known members of this family (Fig. 3) that is also called the pfkb family in the Prosite collection of sequence patterns (Bairoch, 1992). The first eukaryotic member of the family was recently identified in yeast (Thierry et al., 1992). Multiple alignment, identification of conserved regions, and a subsequent database search with corresponding property patterns (Bork & Grunwald, 1990) allowed us to recognize an additional member of this family, inosine-guanosine kinase from *E. coli* (Table 1; Fig. 3), which also is a sugar kinase in that it phosphorylates the ribose of inosine or guanosine.

Divergence of specificity before species divergence

The evolutionary tree (Fig. 4) based on the multiple sequence alignment (Fig. 3) suggests that in this family divergence of specificity occurred before species divergence: fructokinases branched off from ribokinases (Fig. 4, mid-

dle and right) before the divergence of yeast and *E. coli* (Fig. 4, right). That substrate specificity is the major organizing principle of the tree is also evident from the fact that phosphosugar kinases form a major cluster (Fig. 4, left), with early divergence of the approximately equidistant subfamilies 1-phosphofructokinases (K1pf), minor 6-phosphofructokinases (K6p2), and 6-phosphotagatokinases (K6pt). Apparently, the changes required to change specificity from the 1 to the 6 hydroxyl position of phosphofructose are about as difficult as the change in specificity from phosphofructose to phosphotagatose.

The galactokinase family

The third structurally distinct family (Fig. 5) is characterized by three highly conserved boxes (Adams et al., 1988). The sequence segments between these boxes are highly variable in length and amino acid composition. We constructed consensus patterns for the three conserved regions and clearly identified rat mevalonate kinase (Tanaka et al., 1990), its yeast equivalent (RAR1; Kearsley & Edwards, 1987; Oulmouden & Karst, 1991) and phosphomevalonate kinase (EC 2.7.4.2) as members of this family. The regions most conserved between the mevalonate kinases are exactly the conserved boxes of the galactokinases (Schafer et al., 1992). The second box is particularly strongly conserved. A further search with a sequence pattern for only the second box reveals a closely related sequence box in homoserine kinases (EC 2.7.1.39). This box is the most conserved of five conserved regions in homoserine kinases (Mannhaupt et al., 1990). None of the other four match any of the remaining patterns of galactokinase. These facts suggest a common function for the single conserved segment of galactose, mevalonate, phosphomevalonate, and homoserine kinases (e.g., an ATP binding site, as proposed by Tsai & Robinson [1991]).

Jumping into another pathway?

Mevalonate kinase, phosphomevalonate kinase, and homoserine kinase are examples of enzymes homologous to sugar kinases but located in different metabolic pathways. This homology raises an interesting evolutionary

K6pt/Lacla	35	AGGRGLNVT...RVLYESGDKVTATGFLGGK	-152-	GIEWIVVSLGRNGAFAPAK	-11-	DIPVNVVPGSGDSTVAGIASAL
K6pt/Strmu	35	PGGRGLNVT...RVLSQLGDDVSLASGLLGGK	-152-	GIDWIVVSLGSGQAFAPAK	-11-	KIAVNVVPGSGDSTVAGITSAL
K6pt/Staau	35	AGGRGLNVT...RVLAQVGEVPLASGPIGGE	-152-	GIEWIVVSLGAGQAFAPAK	-11-	TISVNLVPGSGDSTVAGITSAL
K1pf_Xanep	38	AGGRGINVA...ACLADWGSQVAALGVLVGV	-154-	GIQLVVISMGTGALFV	-11-	RLAQSSVVGAGDAMVAGLAAAL
K1pf_Rhoca	38	AGGRGVNVA...SFLAHVGHGVAVTGLLGAE	-151-	GVGLVAVSMGAGGAVLV	-11-	ATPIASTVAGDAMVAGLIHAA
K1pf_Ecoli	38	AAGRGINVA...KVLKDLGIDVTVGGFLGKD	-149-	GIAHVVISLGAEGALVW	-11-	SVDVNSTVGAGDSMVGGLIYGL
K6p2_Ecoli	37	PGG.GINVA...RAIAHLGGSATAIFPAGGA	-150-	KAKRVVSLGPPQALGV	-11-	ALKSQSTVAGDRLVGMATLKL
Scrk/Klepn	27	PGGAPANVA...VGVARLGGDSGPIGRVGGD	-155-	QPTELLVLTQKAGVQAA	-11-	PVVAVDPTGAGDAFVAGLLAGL
Scrk_Salty	27	PGGAPANVA...VGVARLGGNSGPIGAVGGD	-155-	QPELLVLTQKAGVQAA	-11-	PVASVDPTGAGDAFVAGLLASL
Scrk_Vibal	26	PGGAPANVA...VAIARLSGKSAPFFGRVGGD	-154-	NIALVLTQKAGVWRV	-11-	VVSPIDTTGAGDAFVGGLLACL
Rbsk_Ecoli	40	PGGRGANQA...VAAGRSGANIAPACTGDD	-147-	GIRTVLITLGRGVWAS	-11-	RVQAVDTIAAGDTFNGALITAL
Rbsk_Yeast	36	AGGRGLNQA...AAIGKLNKPNSSRYSVRMIG	-176-	KRGIVVMTLGRGVLFV	-14-	NVSVDPTGAGDTFLGGLVLTQL
Ingk_Ecoli	92	AGGTIGNTMHNSVLAADDRSVLLGVMCNSIE	-156-	WVDLVLTAGPIGLYMA	-55-	PEKIMNTNGAGDGAALALLHDI

Fig. 3. Ribokinase family: aligned conserved regions. Each of the three sequence "boxes" alone is sufficient to identify the members of the family detected so far. Substrate specificities and protein names are as in Table 1.

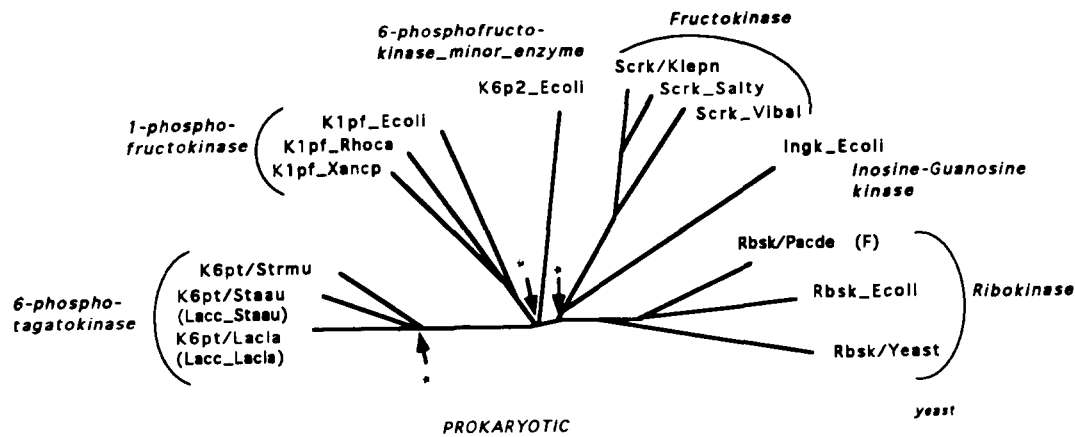


Fig. 4. Ribokinase family: evolutionary tree. Calculated with the program Clustal V using full-length sequences. Fructokinases in this family are structurally distinct from those in the hexokinase family (Fig. 2), a result of independent (convergent) evolution. Notation of protein names as in Table 1.

question. Homoserine kinase catalyzes an important step in the biosynthesis of the amino acid threonine; all other enzymes of this pathway are pyridoxal phosphate-dependent, homologous to one another, and have apparently evolved via several gene duplication events (Bork & Rohde, 1990). So how does homoserine kinase fit in? Perhaps the energy-consuming ATP-dependent homoserine kinase has been inserted into the pathway during evolution to overcome an energy barrier in threonine synthesis. Another puzzling observation is that mevalonate kinase is apparently less similar in sequence to phosphomevalonate kinase (related substrates in subsequent steps of the cholesterol synthesis pathway) than it is to galactokinase (sugar substrate in a completely different pathway).

The galactokinase family, like the other two major families of sugar kinases, has a wide diversity of substrate specificity, all accommodated within the same structural framework, in spite of the fact that the chemistry of the substrates for galactokinase and mevalonate kinase is

quite different (phosphorylation of the first versus the last hydroxyl group).

Conclusions

Three distinct families of sugar kinases

We have compared more than 60 sequences of sugar kinases with 20 different sugar-binding activities. Three major families can be distinguished, here called the hexokinase, ribokinase, and galactokinase families, corresponding to three biologically abundant sugars. There are strong reasons to believe that these three sugar kinase families have different three-dimensional structures: (1) there is no significant sequence similarity between the three families; (2) quality, length, and distribution of the conserved boxes within each family are different; and (3) secondary structure predictions result in very different orders and frequencies of alpha-helices and beta-strands (data not shown).

GAL1_ECOLI galactose	19	THTIQAFGRVNLIGEHTDYNDGFVLP	CAID	62-	GVDVMSGNVPQAGLSSSASLEVA	VGTV	-195-	KGGVMTGGGFGGCI	VALIPEEL			
GAL1_SACCA	44	DFVARS PGRVNLIGEHDYCDFSVLP	LAID	78-	GLQVFCGDIPTGSLSSAAFI	CAVALA	-277-	SYGSRLTGAGWGGCT	VHLVPGGP			
GAL1_KLULA	35	FFITRSPGRVNLIGEHDYCFHV	PMASE	75-	GMEIYVKGDI	PSGGGLSSAAFI	CAVSLA	-275-	SFGSRLTGAGWGGCT	VHLCSTDT		
GAL1/LACHE	18	KDVFPSPGRIN	VIGEHTDYNGGHVFP	PAPIS	66-	GPNLYIEANL	PSGSLSSAAIEM	LMGII	-192-	VLGARMIGGFGGSA	IAIVKKSE	
GAL1_STRLI	22	RGCGRRAGREN	LIGEHTDYNDGFVMP	SPCR	60-	GADVHLASTV	PSGAGLSSAALEVR	PLAM	-195-	GPRRRMTGGGFGGSA	IVLVEAAA	
GAL3_YEAST gal3 protein	39	ILSLGLLGRVNLIGEHDYCDFSVLP	LAID	78-	GAQIFCQSDIPTGGGLSSAFT	CAGRLATI	*					
KIME_RAT mevalonate	5	VLLVSAPGKVLHGEHAVVHGK	VALVALN	93-	SLDIMVWSEL	PPGAGLSSAAYS	VCVAAA	-169-	GLHSKLTGAGGGCG	ITLLKPLG		
KIME_YEAST	4	PFLTSAPGVII	FGESAVYKNPA	VASVS	97-	NIKFSLKSTL	IGAGLGSSAS	ISVSLALA	-184-	IGSTKLTGAGGGCS	LTLLRRDI	
ERG8_YEAST P-mevalonate	4	LRAPSAPGKALLAGGYVLD	TKYEAFV	VGLS	-108-	SPHSHRIE	EVV	.KTGLGSSAGL	VTVLT	-322-	VLTCLIPGAGGYDA	IAVLTQDV
KHSE_BACSU homoserine				80	PVHVKVS	DIPLARGLGSSAAA	IVAAI	EEL				
KHSE_BRELA				85	GLRVVCH	NNIPQSRGLGSSAAA	AAVAG	VAA				
KHSE_CORGL				85	GLRVVCH	NNIPQSRGLGSSAAA	AAVAG	VAA				
KHSE_ECOLI				81	PVAMTLE	KNMFIGSGLGSSAC	SVVAAL	MA				
KHSE_FREDI				82	SVKIEI	DLVPLARGLGSSATA	IVGGL	VLA				
KHSE_YEAST				92	GTKVHV	SNPIPLGRGLGSSGA	AAVAG	VVL				

Fig. 5. Galactokinase family: aligned conserved regions in enzymes structurally related to galactokinases. The pattern describing the central box is the only one in common with homoserine kinase and is the most stringent and discriminating pattern (the random background of unrelated enzymes is separated from the true positive sequences by more than eight mismatches, see Materials and methods). Homoserine kinase fits the pattern with only one mismatch and therefore is proposed to belong to the family. For interpretation see text. Substrate specificities and protein names are as in Table 1.

Why different structural families?

If sugar kinase activity with a wide range of specificities can be constructed on different structural frameworks, then why do we see at least three distinct structural families? Are there subtle ways in which one particular structure performs the function of sugar phosphorylation differently? To answer these questions definitively we need much more biochemical information about each of the enzymes: details of the catalytic mechanism, rate constants, quantitative specificities, and regulatory dependencies. Lacking such information, we can only state some possible reasons. Perhaps at a fairly early stage in evolution, the three classes reflected different basic metabolic roles of different basic sugars, such as glucose as a source of energy or ribose as a central raw material in the pentose phosphate cycle and in nucleic acid biosynthesis. Alternatively, different folds are perhaps needed for different types of metabolic regulation. There is another possibility—nature may simply have reinvented kinase activity on different occasions as a result of historical accident, exploiting different protein folds available at that time in that organism when one basic fold would have done just as well. Whatever happened early in evolution, the three frameworks have, by now, adapted to very diverse specificities in different organisms.

Convergent evolution: Independent evolution of similar specificity

In addition to the surprisingly broad distribution of specificities, the evolutionary trees of the hexokinase and ribokinase families provide multiple evidence of convergent evolution in the sense that similar specificity evolved independently in different structural families (examples: fructokinases) or in different branches of the same structural family (example: glucokinase). Two specific examples follow.

Fructokinases

Flexible adaptation of different structural frameworks to the ability of phosphorylating a particular sugar is evident for fructokinases. The *Zymomonas* fructokinase (Frk/Zymmo; Fig. 2, middle; Zembrzuski et al., 1992) belongs to the hexokinase family, has all of the characteristic sequence patterns of this family, and can be significantly aligned with other members of this family, e.g., it has 26% amino acid identity over 300 residues with *Streptomyces* glucokinase. The other fructokinases are members of the ribokinase family (Fig. 3; Fig. 4, middle), i.e., they are built onto a different three-dimensional structure.

Glucokinases

Also, within one structural family we observe independent adaptation to a similar specificity. For example,

the ability to specifically select a glucokinase substrate has arisen at least twice in the hexokinase family, apparently independently. Yeast and mammalian glucokinases (Hxkg_Yeast, Hxkh_Rat) are in the left branch of Figure 2, grouped with hexokinases, whereas the bacterial glucokinases Hxkg_Zymmo, are in a separate branch (Fig. 2, middle), grouped with fructokinases. Apparently independent evolution of similar specificity (substrate: adrenaline) has also been observed in different branches of the evolutionary tree of G-protein-coupled receptors (Doolittle, 1991). We use the term convergent evolution to describe the independent evolution of the same specificity, the same or different structural scaffolds, as well as the independent evolution of similar catalytic function on different structural frameworks—without implying that details of active site geometry are necessarily similar.

Gene duplication, not exon shuffling

What are the mechanisms for generating so much diversity so easily and occurring so many times? We cannot entirely exclude some kind of domain shuffling mechanism, but at least for the hexokinase family, where we know the three-dimensional structure, there is no obvious compact structural module that carries sugar specificity, nor have we so far seen any sequence evidence for domain shuffling. We do have some evidence that genomic organization (data not shown) appears to be uniform in one family and different between the different structural families. In several cases, a set of enzymes belonging to the same metabolic pathway are coded on one operon. In these cases the entire operon appears to have been copied as a unit before specificity diverged or converged—a very simple and the most plausible mechanism.

Convergent evolution: Other examples

Evolution of similar enzymatic function on different structural frameworks is not an entirely uncommon event (Fig. 6A). A classical example is that of serine proteases: the Ser-His-Asp triad is present in an almost identical three-dimensional constellation in the distinctly different structural frames of trypsin and subtilisin (and their relatives) (Wright et al., 1969). Two more recent examples: there are at least two structurally distinct classes of tRNA synthases (Nagel & Doolittle, 1991) and two different structural classes of superoxide dismutases (Smith & Doolittle, 1992). On several occasions nature appears to have reinvented similar enzymes and even similar specificity using distinctly different three-dimensional folds.

Flexible adaptation of protein structures to diverse functions

These examples of convergent evolution support the idea that active sites can be adapted quite easily, whereas three-

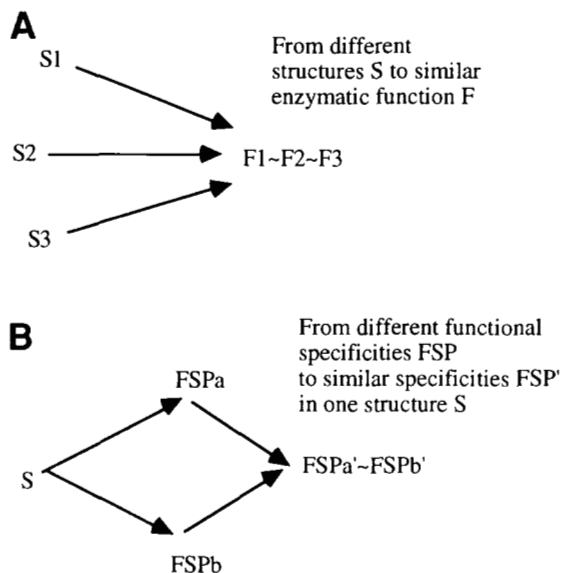


Fig. 6. Two kinds of convergent evolution. **A:** Independent evolution of functionally equivalent active sites in each of three different three-dimensional frameworks. **B:** Independent evolution of the same substrate specificity in different branches of one structural family.

dimensional folds are extremely persistent in evolution. Not only does this appear to be a basic principle of the natural evolution of enzymes, but it is of practical value in protein engineering. In designing and optimizing enzymatic function, the protein engineer can, following nature, rather freely combine different protein structural scaffolds with a variety of specific active site architectures.

In summary, sugar kinases can be grouped by their sequence similarity (and probably by their three-dimensional structures) into three distinct families. Convergent evolution has resulted in functionally equivalent active sites in each of the three different frameworks (Fig. 6A). Within the structurally related families the substrate specificity can be modified without affecting the basic protein fold (Fig. 6B).

Materials and methods

Sequence data

Sequences were taken from the SWISSPROT protein sequence database (release 21; Bairoch & Boeckmann, 1992) or from the literature. All the identifiers are given in Table 1. Sequences from sources other than SWISSPROT are: rat hexokinases type II (Hxk2/Rat; Thelen & Wilson, 1991) and type III (Hxk3/Rat; Schwab & Wilson, 1991), human liver glucokinases (Hxkh/Human; Tanizawa et al., 1991), mouse pituitary glucokinase (Hxkp/Mouse; Hughes et al., 1991), a fragment of a putative isoenzyme of human brain hexokinase (Hxst/Human; Adams et al., 1992), liver cattle hexokinase (Hxkc/Bovin; EMBL accession number

M65140; partly published in Griffin et al., 1989), glucokinase from *Streptomyces coelicolor* (Hxkg/Strc; EMBL accession number X65932; S. Angell, E. Schwarz, & M.J. Bibb, unpubl.), rhamnokinase from *Salmonella typhimurium* (Rhak/Salty; Nishitani & Wilcox, 1991), xylose kinases (Xylk/Stru from *Streptomyces rubiginosus*, EMBL accession number M73789, H.C. Wong et al., unpubl.; Xylk/Artba from *Artherobacter*, Loviny-Anderton et al., 1991; Xylk/Lacba from *Lactobacillus brevis*, Bor et al., 1992; Xylk/Therm from *Thermus thermophilus*, Dekker et al., 1991), yeast ribokinase (Rbsk/Yeast; Thierry et al., 1992), fructokinases from *Klebsiella pneumoniae* (Srk/Klep; Aulkmeyer et al., 1991) and *Zymomonas mobilis* (Frk/Zymo; Zembruski et al., 1992), 6-phosphotagatokinase from *Streptococcus mutans* (K6pt/Strmu; EMBL accession number M80797; Rosey & Steward, 1992), and galactokinase from *Lactobacillus helveticus* (Gal1/Lache; Mollet & Pilloud, 1991).

Property pattern searches in sequence databases

In order to identify additional members of each family, property pattern searches (Bork & Grunwald, 1990) were carried out as follows: The most conserved regions were extracted from a multiple sequence alignment and to each position of these regions a set of steric and physicochemical amino acid properties were assigned. The amino acid property pattern was then translated to a consensus pattern of preferred amino acids, providing a generalization of the original observed amino acid frequencies at each position, and the database of known sequences was screened for similarities. To identify remotely related proteins, deviations from the consensus pattern (mismatches) were allowed. A mismatch is a deviating amino acid property relative to the pattern. In this way a single amino acid difference can be translated to more than one mismatch depending on its physicochemical relation with the other amino acids present in this position. The overall mismatch score is calculated as the sum over all positions of the pattern. If the mismatch score for a particular sequence is well below the random background of scores of apparently nonrelated proteins, this is interpreted as a sequence homology, and the sequence is included in the family for the next iteration of multiple sequence alignment, pattern definition, and database search. Depending on the permissiveness of the cutoffs used, this procedure either explodes by including very many unrelated proteins, not well separated from background, or it converges to a situation where the scores for all family members are well separated from the background. Convergence after a few iterations, with minor adjustments of the multiple sequence alignment, is taken as a strong indication of significance. We use this empirical rule, as a mathematically sound statistical significance estimate has yet to be derived for pattern searches based on multiple sequence alignment insertions and deletions.

Other sequence alignment tools

As an additional, independent check, all sequences included in a family were individually compared with the protein sequence database using the program Fasta (Pearson & Lipman, 1988). Multiple sequence alignments were carried out using MaxHom (Sander & Schneider, 1991) or Clustal V (Higgins et al., 1992). Tree calculations were performed by the neighbor-joining method of (Saitou & Nei, 1987) using Kimura's correction for multiple hits (Kimura, 1983) as implemented in Clustal V (Higgins et al., 1992). Accordingly, insertions/deletions were not considered in calculation of sequence distances. The reliability of the sequence distances represented by the length of tree branches was estimated with a bootstrapping method included in the Clustal V package. The bootstrapping method assesses the statistical reliability by generating a large set (5,000 in this case) of independent trees, each of them representing a different random selection of the sequence positions from the multiple sequence alignment. Thus, the fewer the positions of the multiple sequence alignment that support a grouping the lower is the reliability of the corresponding branch.

Note added in proof

After acceptance of our manuscript, the sequence of a human galactokinase with similarity to the respective family described here was published (Lee et al., 1992). This sequence contains the consensus patterns and underlines the widespread occurrence of this family of sugar kinases.

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