## Research

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# The ABC transporter gene family of *Caenorhabditis elegans* has implications for the evolutionary dynamics of multidrug resistance in eukaryotes Jonathan A Sheps<sup>\*</sup>, Steven Ralph<sup>\*\*</sup>, Zhongying Zhao<sup>+</sup>, David L Baillie<sup>+</sup> and

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## Abstract

**Background:** Many drugs of natural origin are hydrophobic and can pass through cell membranes. Hydrophobic molecules must be susceptible to active efflux systems if they are to be maintained at lower concentrations in cells than in their environment. Multi-drug resistance (MDR), often mediated by intrinsic membrane proteins that couple energy to drug efflux, provides this function. All eukaryotic genomes encode several gene families capable of encoding MDR functions, among which the ABC transporters are the largest. The number of candidate MDR genes means that study of the drug-resistance properties of an organism cannot be effectively carried out without taking a genomic perspective.

**Results:** We have annotated sequences for all 60 ABC transporters from the *Caenorhabditis elegans* genome, and performed a phylogenetic analysis of these along with the 49 human, 30 yeast, and 57 fly ABC transporters currently available in GenBank. Classification according to a unified nomenclature is presented. Comparison between genomes reveals much gene duplication and loss, and surprisingly little orthology among analogous genes. Proteins capable of conferring MDR are found in several distinct subfamilies and are likely to have arisen independently multiple times.

**Conclusions:** ABC transporter evolution fits a pattern expected from a process termed 'dynamiccoherence'. This is an unusual result for such a highly conserved gene family as this one, present in all domains of cellular life. Mechanistically, this may result from the broad substrate specificity of some ABC proteins, which both reduces selection against gene loss, and leads to the facile sorting of functions among paralogs following gene duplication.

## Background

ATP-binding cassette (ABC) transporters are one of the largest families of transport proteins constituting the single largest gene family, comprising about 5% of the genome, in *Escherichia coli* [1]. ABC transporters are grouped into several structural classes, or subfamilies, on the basis of amino

acid sequence and domain organization [2] (Figure 1). The presence of a strongly conserved ATP-binding motif defines membership in the family and the basic functional organization of an ABC transporter in the membrane is the same from bacteria to humans, and in all subclasses [3-5]. A complex of at least two ATP-binding domains, coupled to two blocks of membrane-spanning helices, appears to be the minimum requirement for a functional transporter. Often these domains are found in tandem within a single molecule, but in many cases are distributed across separate proteins that must then assemble in the membrane. ABC transporters are collectively able to accommodate an unusually large array of different substrates. This diversity of function is manifest at the family level, but also in individual members of the family, for example those associated with multidrug resistance (MDR).

Decottignies and Goffeau [6] catalogued the entire ABC transporter family of the yeast *Saccharomyces cerevisiae* and in so doing delineated six of the major subgroups of eukaryotic ABC transporters. Allikmets *et al.* [7] catalogued all the then known 33 human ABC transporters, including those known only from partial expressed sequence tag (EST) sequences, and divided these into seven subfamilies. This scheme has been adopted, with a revised nomenclature, by the Human Genome Organisation (HUGO) [8] in order to provide a unified nomenclature for both human and mouse ABC transporters. Of these seven subfamilies, one, ABCA, has no exact equivalent in the yeast genome [9,10]. Genes considered to be part of subfamily ABCA have been identified in the slime mold *Dictyostelium discoideum*, as well as in malaria



#### Figure I

Structural diversity of ABC transporters. Illustration of the various domain organizations found among members of the ABC transporter family in *C. elegans.* TM indicates a transmembrane domain typically containing six predicted membrane-spanning helices. ABC indicates an ATP-binding cassette domain. The color codes for each structure are used throughout the figures to show the lack of concordance between structural categories and families defined on the basis of sequence similarity.

parasites [11] and *Caenorhabditis elegans* (this paper). With the completion of the human and *Drosophila melanogaster* genomes, a joint summary of the ABC transporter complements of both genomes was published [12]. This identified a new subfamily, ABCH, which appears to be the most divergent yet. One, previously unclassified yeast ABC gene, YDR061w [13], appears to be a structurally aberrant member of subfamily H.

The phenotypes of five ABC transporter knockouts have been reported in C. elegans. Four of these involve genes expected, by homology to mammalian genes, to be involved in drug resistance: three P-glycoproteins (Pgp-1, Pgp-3 and Pgp-4) (subfamily B) and one multi-drug resistance protein (MRP) [14,15] (subfamily C). These ABC transporter mutants are associated with sensitivity to environmental insult [16]. Pap-3 mutant strains of C. elegans are more sensitive to the drugs chloroquine and colchicine. Pgp-1 and mrp-1 strains are hypersensitive to toxic pigments produced by some bacteria [17]. All the nematode P-glycoproteins examined so far seem to be highly expressed in intestinal cells [18], and in the excretory cell, which functions somewhat like a kidney in C. elegans. The mrp-1, pgp-1 and pgp-3 mutant strains have been reported to be hypersensitive to the heavy metals cadmium and arsenite [15]. The fifth reported knockout is of the product of the ced-7 gene [19]. Mutant alleles of ced-7 cause a defect in engulfment of the cell corpses left behind by apoptosis. ced-7 is a member of the ABCA subfamily, and has a similar phenotype to the abca1 gene in humans. ABCA1 protein is required for engulfment of apoptotic cells by macrophages and is thought to regulate membrane fluidity through an increase in phosphatidylserine exposure on the outer leaflet of the cell membrane [20].

The term orthology is used to describe genes separated from one another by speciation events while paralogy describes those separated by gene duplication events [21]. Of particular interest, from the point of view of functional annotation, are the cases where a pair of genes, one from each of a pair of organisms, are found. In these cases it is reasonable to presume that the orthologous genes may share a conserved function retained from the same single gene present in the common ancestor of the two organisms. However, where a single gene (or set of duplicated genes) in one genome is most closely related to a set of duplicated (paralogous) genes in another genome this is sometimes termed co-orthology [22], and then no particular orthologous pair can be unambiguously specified. In the case of co-orthologs the argument for retention of analogous functions between members of the sets of descendant genes is much weaker. Comparison of two complete genomes, those of C. elegans and S. cerevisiae [23], demonstrated a high fraction of ortholog pairs in gene families involved in core biological functions. Specifically, Chervitz et al. [23] found, when pairing conserved yeast genes with their most similar worm homologs (subject to a BLAST score cut-off of  $< 10^{-100}$ ), 57% of these highly conserved gene pairs

involved orthologous, rather than paralogous, pairs of genes. In this category of core functions they included trafficking, and, as possibly the largest family of trafficking genes in animal genomes, ABC transporters should be expected to share in this high level of one-to-one correspondence between genomes. We expected therefore that this would allow us to assign predicted functions to newly discovered C. elegans ABC proteins on the basis of their already-characterized mammalian orthologs. Following a comprehensive phylogenetic analysis of ABC transporters from four eukaryote genomes, we found that the frequency of orthologous pairs among ABC transporters was substantially lower than we expected. Particular domain organizations and substrate specificities seem to have evolved independently several times in multiple lineages. This is expected to complicate the functional analysis of ABC transporter function in newly characterized genomes.

## **Results and discussion**

Here we present a classification of all ABC transporters encoded in the C. elegans genome, based on a phylogenetic analysis which includes the 49 currently known human ABC proteins for which there are reliable, public, sequence data. We took the approach of analyzing primarily the conserved ATP-binding cassettes from each protein, regardless of the structural class from which the domain is drawn. This allows evaluation of the evolutionary history of each protein in the family, without biases that might result from gene-fusion events resulting in convergent acquisition of similar domain structures by distantly related proteins. In addition, we reevaluated the relationships of transporters within statistically reliable clusters whose members are closely related enough that structural variations do not lead to errors in alignment. We did this to capture additional phylogenetic information, which may be apparent in the less conservative transmembrane domains, at a level of analysis where it is less likely to be misleading.

An example of our first-pass approach is given in Figure 2, which shows an analysis of isolated ATP-binding cassette domains from the human ABC transporters only. In particular, we find that all seven subfamilies recognized by Allikmets *et al.* [7] are recovered with significant bootstrap support. Their finding, that subfamily B is more closely related to the carboxy-terminal component of subfamily C than the two halves of ABCC molecules are with one another, is supported by our results.

## A collection of transporters

We found a total of 60 confirmed ABC transporters in the annotated protein set derived from the *C. elegans* genome sequence. This represents approximately 0.3% of the total number of genes (approximately 19,000) in the worm genome. Only 8 of the 60 predicted genes lack any corresponding mRNA (Table 1), and only one (F56F4.6) is

structurally aberrant in a way that would suggest it is likely to be a pseudogene.

Thirty ABC transporters are described in the yeast genome, or approximately 0.5% of its approximately 6,000 proteins [13]. At present 49 human ABC transporters have been identified and, at least partially, cloned. They are included here (Figures 3,4,5,6,7 and Table 2) to illustrate their relationships with nematode proteins, which might then shed light on their biological roles. Inclusion of human as well as *D. melanogaster* ABC transporters in our tree allows us to explicitly classify *C. elegans* ABC transporters according to the current eight-subfamily taxonomic scheme for ABC transporters [12].

## Typing ABCs to subfamily

We define membership of a particular gene in an ABC transporter subfamily primarily on the basis of the position of its ATP-binding domains in our first phylogenetic tree (not shown). Genes that fell unambiguously within a clade containing genes already assigned to given subfamily, were included in that subfamily. Where we could not assign a gene to a particular clade with a significant bootstrap value, the assignment was made on the basis of which subfamily's members scored highest when that gene was used as query in a BLAST search. The subfamilies are sometimes named according to the well-characterized mammalian genes that typify each of them, for example, P-gp (P-glycorprotein), MRP, White gene homologs, RNAse L inhibitor, GCN20 homologs, ABC1 and ALDP [7]. These correspond to the HUGO-defined subfamilies B, C, G, E, F, A and D, respectively. Re-analysis of the full-length sequences confirmed the placement all C. elegans genes within the preexisting subfamilies, with substantial bootstrap support (Figures 3,4,5,6,7).

## Instances of orthology

In the set of worm and human ABC transporters, only 8 of 49 possible pairs (16%) of sister genes contained a single human protein and a nematode homolog (Table 3). Similarly, 10% of ABC transporters were found in orthologous pairs when the comparison is made between yeast and worm genomes. A more comprehensive comparison of worm and yeast genomes [23] came to the overall conclusion that 57% of genes in highly conserved gene families were found in orthologous pairs, and the study suggested that such gene families provide a conserved core proteome which forms the basis of eukaryote biochemistry. ABC transporters are conserved in all eukaryotic and prokaryotic genomes, so it is interesting to note that they are found in orthologous pairs much less frequently than most gene families that are roughly as well conserved. Clearly, ABC transporter evolution has not been typical of strongly conserved gene families, and while we might have inferred that ABC-transporter-mediated metabolism differs radically among eukaryotes, this seems improbable, given the broadly comparable set of substrates associated with ABC transporters in all eukaryotes where they have been studied.

Figure 2 (see legend on next page)



#### Figure 2 (see previous page)

Tree of human ATP-binding cassette domains. The evolution of the ABCB subfamily from within the ABCC subfamily, and the structural diversity of subfamily B is shown here. Each cluster of ABC domains within each subfamily, except for subfamily B, is collapsed to form a single, representative, branch; n-term: amino-terminal ABC; c-term: carboxy-terminal ABC. The phylogeny of ATP-binding cassettes from human ABC transporters was produced according the following procedure. Predicted amino-acid sequences were aligned using ClustalX [54]. Aligned sequences were used to generate matrices of mean distances among proteins, and these matrices were used to generate a phylogenetic tree according to the neighbor-joining algorithm [55], refined using the SPR branch-swapping technique under the minimum evolution criterion, implemented by PAUP\*4.0b10 [56]. Bootstrapping [57] was used to determine the relative support for the various branches of the tree (1,000 replicates), and nodes with less than 50% support were collapsed to form polytomies. The structures of the proteins in which the domains are embedded are indicated according to the color scheme in Figure 1. It should be noted that branch lengths in the figures are not to scale and do not represent distances between protein sequences. The original alignment files are available as Additional data files 1-8.

Within the P-gp-related ABCB subfamily, the only one-to-one pairings found between *C. elegans* and human genes are those of Wo9D6.6 (Haf-5) and MTABC3 (B6), and Y48G8AL.11 (Haf-6) and MABC1 (B8). These are both half-transporters localized to the mitochondria. MTABC3 (B6) is involved in iron homeostasis [24] and its rat ortholog, PRP, is overexpressed during hepatocarcinogenesis [25]. Two other mitochondrial ABC transporters in humans, MABC2 (B10) and ABCB7, have orthologs in flies and/or yeast, but not nematodes.

Among ABCC molecules, whose range of functions broadly overlaps with P-gps, only C18C4.2 (Cft-1) and CFTR (C7) are indicated as orthologs in our analysis. However, the bootstrap value on this pairing is very low (51%, see Figure 5), so we cannot attach much confidence to this observation. It may simply be that C18C4.2 (Cft-1) is a highly divergent member of subfamily C, and does not bear much functional similarity to CFTR (C7). Although not forming simple pairs with any nematode gene, human MRP5 (C5), a transporter of nucleotide analogs [26,27], and ABCC11 and ABCC12 appear to be coorthologous to worm F14F4.3 (Mrp-5), which may provide some hint as to the function of the latter.

All four of the C. elegans members of subfamilies E and F (Figure 6) form strongly supported and unambiguous pairs with their homologs in D. melanogaster, Homo sapiens, and yeast. This unusually strong conservation, compared to the other subfamilies of ABC genes, argues for involvement in something indispensable, at least on an evolutionary timescale. The three genes in subfamily F, which lack transmembrane domains, are generally regarded as forming ribosome associated proteins involved in regulation of mRNA translation, rather than transporters. The RNase L inhibitor (E1), also known as the oligoadenylate-binding protein (OABP), is thought to be involved in the regulation of the interferoninduced antiviral response [28] that bears some similarities to the mechanism thought to underlie the now common molecular biology technique of double-stranded RNAdirected interference (RNAi). It also seems to have a role in muscle differentiation [29] in mammals. The critical role of the RNase L inhibitor is underlined by its conservation even in a highly reduced genome. In the rather minimal genome of the endosymbiotic Guillardia theta nucleomorph (302 genes)

the RNase L inhibitor is the only ABC protein found [30]. The yeast ortholog of the RNase L inhibitor protein, YDR091c, is essential for growth, as is YER036c, the yeast ortholog of T27E9.7/ABCF2 [31]. On the other hand GCN20, the yeast version of F42A10.1/ABCF3, is not essential, although mutants do have specific defects in translation.

## Processes of gene duplication and loss

While the conservation of simple orthologous gene pairs is a rare observation in our study, the numbers of genes in most ABC transporter subfamilies are about the same, despite numerous instances of gene duplication and loss. For example, within ABCB the number of half-transporters in each genome is almost identical. Furthermore, most mammalian half-transporters in subfamily B are found in clusters of functionally related, or at least co-localized, genes (the TAP (B2 and B3) genes, and the four mitochondrial ABCB genes, MABCs1 and 2 (B8 and B10), MTABC3 (B6) and ABCB7 [32]), paired with similarly sized groups of C. elegans genes. Likewise the number of genes in subfamilies A, C and D is much the same between genomes. However, it does appear that C. elegans, relative to humans, has undergone a massive expansion in the P-gp (full or pseudo-dimer configuration) subclass of subfamily B, and subfamily G, the 'White-like' genes. The likelihood that ABC transporter lineages have been lost repeatedly in evolution is evident from the phylogeny. The single group of P-gps in mammals contains only four members, while *C. elegans* has 15 P-gps, of which only three are closely related to their mammalian homologs. A literal reading of the tree (Figure 4) would suggest the presence of five additional P-gp lineages in the common ancestor of nematodes, flies and mammals that have been lost, independently, in both mammals and flies. These losses, and the species-specific expansion of the remaining lineages of genes, underlines the peculiarly dynamic composition of this group of multifunctional transport proteins.

## Conclusions

The completion of the *C. elegans* and *D. melanogaster* genome projects [33,34] make it possible to analyze entire gene families in metazoans. The advantage of performing a combined analysis of all known ABC proteins from two organisms is that it allows unambiguous identification of

#### Table I

## Characterization of the 60 C. elegans ABC proteins

Subfamily		ORF name/CGC name	Chromosome	GenBank accession number	Size (amino acids)	Predicted topology	cDNA if known	RNAi phenotype
A	Abt	C24F3.5/ Abt-1	IV	CAA18775	1,429	(6TM-ABC)2		None
		C48B4.4/ Ced-7	Ш	NP_499115	1,704	(8TM-ABC)2	Complete	None
		FI2B6.1/ Abt-2	I	AAB54153	1,547	(6TM-ABC)2	Partial	None
		F55G11.9/ Abt-3	IV	CAB05222	1,431	(8/4TM- ABC)2		None
		F56F4.6	I	AAB54203	260	ABC		None
		Y39D8C.1/ Abt-4	V	AAC69223	1,802	(6/8TM- ABC)2	Partial	None
		Y53C10A.9/ Abt-5	I	CAA22142	1,564	(6TM-ABC)2	Partial	None
В	Pgp (full molecules)	C05A9.1/ Pgp-5	Х	CAA94202	1,283	(6TM-ABC)2	Partial	None
		C34G6.4/ Pgp-2	I	AAB52482	1,265	(6TM-ABC)2	Partial	None
		C47A10.1/ Pgp-9	V	CAB03973	1,294	(6TM-ABC)2	Partial	None
		C54D1.1/ Pgp-10	х	AAC48149	1,283	(4TM-ABC)2	Partial	None
		DHII.3/ Pgp-II	П	CAA88940	1,270	(6TM-ABC)2	Partial	None
		F22E10.1/ Pgp-12	х	CAA91799	1,318	(6TM-ABC)2	Partial	None
		F22E10.2/ Pgp-13	х	CAA91800	1,291	(6TM-ABC)2		None
		F22E10.3/ Pgp-14	х	CAA91801	1,327	(6TM-ABC)2	Partial	None
		F22E10.4/ Pgp-15	х	CAA91802	1,270	(6TM-ABC)2		None
		F42E11.1/ Pgp-4	х	CAA91463	1,266	(6TM-ABC)2	Partial	None
		K08E7.9/ Pgp-1	IV	CAB01232	1,321	(6TM-ABC)2	Partial	None
		T21E8.1/ Pgp-6	х	CAA94220	1,225	(6TM-ABC)2	Partial	None
		T21E8.2/ Pgp-7	х	CAA94219	1,269	(6TM-ABC)2		None
		T21E8.3/ Pgp-8	х	CAA94203	1,243	(6TM-ABC)2	Partial	None
		ZK455.7/ Pgp-3	х	CAA91467	1,268	(6TM-ABC)2	Partial	None
	Haf (half molecules)	C30H6.6/ Haf-1	IV	CAB02812	586	4TM-ABC	Partial	None
		F43E2.4/ Haf-2	II	AAC71121	761	8TM-ABC	Partial	None
		F57A10.3/ Haf-3	V	CAB09418	733	6TM-ABC	Partial	None
		W04C9.1/ Haf-4	I	AAC68724	787	8TM-ABC	Complete	Weak embryonic lethality, slow growth
		W09D6.6/ Haf-5	Ш	CAB04947	801	8TM-ABC	Complete	None
		Y48G8AL.11/ Haf-6	I	AAK29911	565	4TM-NBF	Partial	
		Y50E8A.16/ Haf-7	V	CAB60586	807	6TM-ABC	Partial	
		Y57G11C.1/ Haf-8	IV	CAB16503	633	4TM-ABC		None
		ZK484.2/ Haf-9	Ι	AAK39394	815	8TM-ABC	Complete	None
с	Mrp/Cft	CI8C4.2/ Cft-I	۷	AAK52175	1247	(5/6TM- NBF)2	Partial	None
		E03G2.2/ Mrp-3	х	CAA92148	1,398	(6TM-ABC)2	Partial	None
		FI4F4.3/ Mrp-5	х	CAB54225	1,427	(6TM-ABC)2	Partial	Slow growth, Clear
		F20B6.3/ Mrp-6	х	AAA82317	1,396	(6TM-ABC)2	Partial	Egg laying defect
		F21G4.2/ Mrp-4	х	CAB02667	1,573	(10/6TM- ABC)2	Partial	None

Characteriz	ation of the 60 C. elegans ABC p	roteins					
	F57C12.4/ Mrp-2	х	AAB07022	1,525	(10/6TM- ABC)2	Complete	None
	F57C12.5/ Mrp-1	х	AAD31550	1,528	(12/6TM- ABC)2	Complete	None
	Y43F8C.12/ Mrp-7	V	CAA21622	1,119	(12/2TM- ABC)2	Partial	
	Y75B8A.26/ Mrp-8	х	CAA22110	1,144	(4/6TM- ABC)2	Partial	
D	C44B7.8	Ш	AAA68339	665	4TM-ABC	Partial	
	C44B7.9	П	AAA68340	661	4TM-ABC	Partial	None
	C54G10.3	V	CAA99810	660	6TM-ABC	Complete	None
	T02D1.5	IV	CAB0590	734	6TM-ABC	Partial	None
	T10H9.5	۷	AAC19238	598	6TM-ABC	Complete	None
E	Y39E4B.I	Ш	CAB54424	610	ABC-ABC	Partial	Embryonic lethality
F	F18E2.2	V	CAA99835	622	ABC-ABC	Partial	None
	F42A10.1	III	AAA19072	712	ABC-ABC	Partial	None
	T27E9.7/ GCN20-2	III	CAB04880	622	ABC-ABC	Complete	None
G	C05D10.3	Ш	AAA20989	598	ABC-4TM	Partial	None
	C10C6.5	IV	CAB05682	610	ABC-6TM	Partial	None
	C16C10.12	III	CAA86750	610	ABC-4TM	Partial	None
	F02E11.1	II	AAB66050	658	ABC-4TM	Partial	None
	F19B6.4	IV	CAA93461	695	ABC-6TM	Partial	None
	T26A5.I	III	AAC77504	608	ABC-4TM	Partial	None
	Y42G9A.6	III	AAF60554	684	ABC-6TM	Partial	None
	Y47D3A.11	III	CAB57891	547	ABC-6TM	Partial	None
	Y49E10.9	III	CAB11549	454	ABC-4TM		None
н	C56E6.1	II	AAA81093	1,667	ABC-12TM		Larval arrest
	C56E6.5	Ш	AAA81094	595	ABC-6TM	Partial	None

Table I (Continued)

Subfamily names given are according to the HUGO nomenclature (A-H) [33] as well as the CGC (*Caenorhabditis* Genetics Centre [61]) gene names for each subfamily. TM, transmembrane domain, where the number preceding it is the predicted number of membrane spanning helices or the number in the amino-terminal/carboxy-terminal TM domains, respectively. ABC, ATP-binding cassette. The existence of known cDNAs, whether complete or partial, is listed according to information in WormBase release WS112 [62]. RNAi phenotypes of genes on chromosome I are given according to [63], those on chromosomes II, IV, V and X are from [64], those of genes on chromosome III are from [65].

orthologous pairs of genes, as well as allowing the pattern of evolution by a process of gene duplication, lineage sorting, and functional convergence to be explicitly modeled.

Saurin *et al.* [35] surveyed the ABC transporters, considering both eukaryotic and prokaryotic systems, and found that there is a fundamental phylogenetic division among ABC transporters involved in import versus export processes. The importer class of ABCs is found only in prokaryotes, whereas exporters are found in all domains of life [35]. However, that survey, while covering all classes of ABC transporter, was not comprehensive with respect to any of the organisms surveyed. Most recently, Schriml and Dean [10] compared the human ABC family to that of the mouse *Mus musculus*, and found almost perfect identity between the two genomes. We have integrated previous information with the complete inventory of ABC transporters from the genome of the nematode worm *C. elegans*. We find that most of the ABC transporters in the worm can be classified into the existing human transporter taxonomy. We find 60 ABC transporters in the worm genome, representing an overall doubling in size of the ABC transporter family relative to yeast, whose genome



#### Figure 3

Phylogenetic tree of ABCA proteins in three eukaryote genomes. A phylogeny derived and displayed according to the procedure outlined in the legend to Figure 2, except that complete protein sequences were used, not just those of the ATP-binding cassettes. The genome of origin for each protein is indicated by prefixes before each gene name, according the following scheme: Ce, *C. elegans*; Dm, *D. melanogaster*; Hs, *H. sapiens*; Sc, *S. cerevisiae*.

contains one third as many protein-coding genes. No ABC genes were found that could be classified among the bacterial import proteins.

At least three subfamilies of ABC transporter contain members capable of a conferring an MDR phenotype, and transporters from at least two different subfamilies cause

#### Figure 4 (see following page)

Phylogenetic tree of ABCB proteins in four eukaryote genomes. A phylogeny derived and displayed according to the procedure outlined in the legend to Figure 3. Shown here is the division between the half transporters, which are most of the ABCB genes in mammals, and the full-transporters (called P-glycoproteins (P-gps)) that have evolved from them. Four lineages of P-gps (exemplified by genes F22E10.1-4, T21E8.1-3, C47A10.1 and C54D1.1) have been lost in both flies and mammals, and of the two remaining P-gp lineages, one has been lost in each of the fly and human lines of descent. Subsequent duplications within the single remaining P-gp lineage in both flies and mammals have not been sufficient to keep pace with continuing P-gp duplications in the worm genome.



Figure 4 (see legend on previous page)

Phylogenetic tree of ABCC proteins in four eukaryote genomes. A phylogeny derived and displayed according to the procedure outlined in the legend of Figure 3.

MDR in human tumors [36]. A multi-drug transporter is a single protein capable of specifically recognizing several structurally distinct classes of compounds, and which catalyzes their efflux from the cell or sequestration in a subcellular compartment. Proteins of the P-glycoprotein (P-gp) group (ABCB) transport hydrophobic compounds and function in transport of lipids and bile from the liver as well as generally defending the body from toxic natural products





#### Figure 6

Phylogenetic trees of ABCD, ABCE, and ABCF proteins in four eukaryote genomes. Phylogenies derived and displayed according to the procedure outlined in the legend of Figure 3.

in the diet [37]. P-gps are also a component of the blood-brain barrier and function in tolerance of drugs normally minimally toxic to mammals, such as ivermectin [38]. Multi-drug resistance mediated by MRP group (ABCC) proteins depends on a slightly different mechanism. MRPs seem to function by cotransporting toxic compounds with glutathione, or as glutathione conjugates [36]. An MDR phenotype is also associated with some members of the ABCG group of transporters, in both yeast [39] and humans [40]. The MDR phenotype appears to have evolved not just once, but at least three times in the history of ABC transporters. Given the distribution of MDR-causing and non-MDR genes among mammalian Pgps; it seems reasonable to infer that MDR genes may well have arisen more than once among the P-gps themselves. It has been observed [41,42] that the entire ABC transporter family is characterized by a highly adaptable common mechanism for coupling substrate binding to ATP hydrolysis and extrusion. It has been pointed out that, because P-gp



#### Figure 7

Phylogenetic trees of ABCG and ABCH proteins in four eukaryote genomes. Phylogenies derived and displayed according to the procedure outlined in the legend of Figure 3.

## Table 2

## Alphabetic list, by taxon, of protein sequences used in this study

ADPI         NP_009937         171D11.2         AAF4509         C05A9.1         CAA94202         ABCA1         NP_00349           ATMI         NP_01308         Atet         AA75102         C05D10.3         AAA2099         ABCA2         NP_01396           BTTI         NP_013086         Brown         AAF7020         C016C5.5         CAB05682         ABCA3         CAS6395           CAFI6         NP_11664         CG10441         AAF53737         C16C10.2         CAB05682         ABCA5         NP_05123           MDL1         NP_013289         CG10505         AAF45706         C24F3.5         CAA18775         ABCA7         NF32277           PDR10         NP_014737         CG1464         AAF5327         C4467.8         AA85439         ABCA1         NF32277           PDR11         NP_01522         CG11890         AAF5661         C3464.4         CA80373         ABCA1         NP_68974           PDR12         NP_01578         CG1703         AAF5667         C47710.1         CA63037         ABCA1         NP_68974           PDR3         NP_01773         CG1703         AAF5667         C47710.1         CA63037         ABCA1         NP_68974           PDR3         NP_015176         CG1703         <	S. cerevisiae	Accession number	D. melanogaster	Accession number	C. elegans	Accession number	H. sapiens	Accession number
ATMI         NP_010308         Brown         AAF47020         C05010.3         AA20989         BEAC3         NP_01097           BPTI         NP_010608         Brown         AAF47020         C106C.5.         CA805825         ABECA3         CAA55825           CAF16         NP_116624         CG10226         CG10226         AAF450670         C16C10.12         CA80730         ABECA5         NP_051142           GCN20         NP_016644         CG10505         AAF45706         C2474.35         CAA16377         ABECA7         AF323787           MDL1         NP_016973         CG11147         AAF52866         C1487.9         CAA63832         ABECA10         NP_055202           PDR11         NP_012527         CG11897         AAF56866         C1487.9         AAA68339         ABECA10         NP_60591           PDR15         NP_010736         CG1703         AAF49018         C4814.9         AA68130         ABCC12         NP_60591           PXA1         NP_01733         CG1494         AAF5033         C566.1         AA81033         ABCC10         NP_269914           PXA1         NP_01733         CG1738         AAF5376         D2111         AA66030         ABCC11         NP_149163           S56         N	ADPI	NP_009937	171D11.2	AAF45509	C05A9.1	CAA94202	ABCAI	NP_005493
BPT1         NP_013086         Brown         AAF3020         C10Cs5         CAB05602         RAD5560         CAB05602           CAF16         NP_116644         CG10241         AAF5070         C16C10.12         CAB6370         ABCA5         NP_051141           GCN20         NP_116644         CG10411         AAF53717         C18C40.12         CAA1575         ABCA6         NP_255023           MDL1         NP_01533         CG11050         AAF64706         C2473.5         CAA16775         ABCA8         AB020629           PDR10         NP_01577         CG11147         AAF53284         C3466.4         AA6339         ABCA10         NP_05567           PDR11         NP_01577         CG1189         AAF56870         C4740.1         CA03073         ABCA12         NP_06547           PDR12         NP_016976         CG12703         AAF6980         C4610.3         CAA8149         ABCC12         NP_05576           PSA1         NP_015778         CG14707         AAF59803         C5461.3         CAA8109         ABCC12         NP_255022           SNQ2         NP_010316         CG12703         AAF69083         C5461.3         CAA81039         ABCC10         NP_25502           SNQ2         NP_010320         C	ATMI	NP_014030	Atet	AAF51027	C05D10.3	AAA20989	ABCA2	NP_001597
CAF16         NP_116625         CC1026         AAF63070         C16C10.12         CAA86730         ABCA6         NP_91121           GCN20         NP_105280         CG10501         AAF63760         C18C4.2         AAK52175         ABCA7         NP_323023           MDL1         NP_012380         CG10505         CG10505         AAF63760         C247.35         CAA163775         ABCA7         NP_323023           PDR10         NP_019533         CG11169         AAF53641         C3464.4         AA85339         ABCA10         XP_085647           PDR11         NP_012527         CG11897         AAF58669         C447.10         AA643340         ABCA12         NP_689914           PDR15         NP_016476         CG112703         AAF49018         C4881.4         CAA8139         ABCC11         NP_689914           PXA2         NP_012713         CG14709         AAF49069         C5666.1         AA81093         ABCC11         NP_18951           Sta6         NP_012713         CG1738         AAF39373         DH11.3         CAA81991         ABCC11         NP_10326           Sta6         NP_01375         CG1738         AAF39373         D141.3         CA88490         ABCC12         NP_103262           YDR61140	BPTI	NP_013086	Brown	AAF47020	C10C6.5	CAB05682	ABCA3	CAA65825
GCN20         NP.116644         CG10411         AAF3737         C18C4.2         AAS1275         ABCA         NP.52023           MDL1         NP.01503         CG10505         AAF6364         C24F3.5         CAA18775         ABCA7         A328787           MDL2         NP.01503         CG11067         AAF5364         C3046.6         CA802812         ABCA8         A8020629           PDR10         NP.01252         CG11807         AAF5367         C4487.4         AA53422         ABCA10         XP.028547           PDR11         NP.015267         CG11898         AAF56870         C4487.9         AAA63340         ABCA12         NP.056472           PDR15         NP.010794         CG1703         AAF4908         C48844         CA63144         ABCB7         A8003289           PXA2         NP.01736         CG1470         AAF48087         C566.5         AAA81092         ABCC10         NP.252621           St66         NP.012713         CG1738         AAF48087         C566.5         AAA81092         ABCC10         NP.252621           St67         NP.010346         CG1801         AAF80836         F0211.1         AA686050         ABC1         NP.010320           St70         PP.010376         CG1810	CAF16	NP_116625	CG10226	AAF50670	CI6CI0.12	CAA86750	ABCA5	NP_061142
MDL1         NP_013289         CG10050         AAF64706         C2473.5         CAA18775         ABCA7         A328787           MDL2         NP_010533         CG11049         AAF5324         C3016.6         CAB02812         ABCA8         AB02029           PDR10         NP_012522         CG11407         AAF5224         C34G6.4         AA85340         ABCA10         NP_05472           PDR11         NP_012527         CG11897         AAF56867         C4747.01.1         CA803973         ABCA12         NP_056472           PDR15         NP_010746         CG12703         AAF49018         C4884.4         CA803973         ABCA13         NP_056472           PXA1         NP_01773         CG14704         AAF56856         C5401.1         AC48144         ABCB5         AA673470           SK6         NP_012733         CG1494         AAF56838         C54G10.3         CA79910         ABCB5         AA60393         ABCC10         NP_258261           SK6         NP_01024         CG1703         AAF5336         DH11.3         CA89804         ABCC13         NP_74921           YDR1         NP_013052         CG1803         AAF5361         F02E11.1         AA86050         ABCF1         NP_10326           YDR016<	GCN20	NP_116664	CG10441	AAF53737	C18C4.2	AAK52175	ABCA6	NP_525023
MDL2         NP_019053         CG11069         AAF5361         C30H6.6         CAB02812         ABCA8         AB020629           PDR10         NP_01973         CG11147         AAF52284         C34G6.4         AAB52482         ABCA10         NP_50502           PDR11         NP_015267         CG11897         AAF58680         C4487.9         AAA68340         ABCA10         NP_056472           PDR15         NP_01694         CG11898         AAF56860         C4487.9         AAA68340         ABCA13         NP_659472           PDR15         NP_01694         CG1203         AAF49016         C4803973         ABCA13         NP_659474           PDR5         NP_01578         CG1203         AAF49016         C48044         CA203973         ABCC1         NP_259261           SNQ2         NP_010734         CG1738         AAF50837         C5665.5         AAA81092         ABCC11         NP_149163           YCF1         NP_010316         CG1744         AAF51314         E03C2.2         CA38940         ABCC12         NP_149163           YCF1         NP_010316         CG1814         AAF50836         F02E11.1         AAB65050         ABCF1         NP140137           YCF1         NP_010316         CG1814 <td< td=""><td>MDLI</td><td>NP_013289</td><td>CG10505</td><td>AAF46706</td><td>C24F3.5</td><td>CAA18775</td><td>ABCA7</td><td>AF328787</td></td<>	MDLI	NP_013289	CG10505	AAF46706	C24F3.5	CAA18775	ABCA7	AF328787
PDR10         NP_019473         CG11147         AAF52284         C34G4         AAB52482         ABCA9         NP_52502           PDR11         NP_015257         CG111897         AAF58366         C44B7.8         AAA68339         ABCA10         XP_085471           PDR12         NP_01576         CG11898         AAF58670         C44B7.8         AAA68339         ABCA13         NP_689914           PDR5         NP_016776         CG12703         AAF58656         C54D1.1         ACA81344         ABCB7         AB0052891           PXA1         NP_015778         CG1470         AAF59083         C54G10.3         CAA99910         ABCB7         AB0052891           PXA2         NP_012733         CG1718         AAF50836         C54G1.3         CAA99910         ABCC1         NP_149163           SNQ2         NP_0103052         CG17338         AAF50837         C56E5.5         AAA81092         ABCC12         NP_01324           YCF1         NP_0103052         CG1891         AAF50846         F12E6.1         AAB5133         ABC72         NP_005683           YDR061w         NP_010360         CG1891         AAF53767         F1286.1         AAB5133         ABC72         NP_005683           YER30AC         NP_011480	MDL2	NP_015053	CG11069	AAF56361	C30H6.6	CAB02812	ABCA8	AB020629
PDR11         NP_01252         CCI1460         AAF5527         C4487.8         AAA68330         ABCA10         XP_065472           PDR12         NP_015267         CGI1897         AAF56680         C47A10.1         CA68373         ABCA12         NP_05472           PDR15         NP_01676         CGI1703         AAF56050         C47A10.1         CA60373         ABCA13         NP_669714           PDR15         NP_01578         CG1703         AAF4018         C4884.4         CA62344         ABCB5         AA605289           PXA1         NP_015178         CG1703         AAF4069         C5665.1         AAA81092         ABCC10         NP_258261           SNQ2         NP_010271         CG1764         AAF50837         C5665.5         AA81092         ABCC13         NP_149163           YET1         NP_01019         CG1764         AAF50847         F1286.1         AAB513         ABCC12         NP_033226           YDR061W         NP_010136         CG1810         AAF50847         F1286.1         AABC410         AAF3488           YDR071C         NP_010360         CG1814         AAF61877         F1474.3         CA854225         ABCF3         NP_006828           YDR070W         NP_010350         CG1814	PDR10	NP_014973	CGIII47	AAF52284	C34G6.4	AAB52482	ABCA9	NP_525022
PDR12         NP_015267         CG11897         AAF56897         C4487.9         AAA68340         ABCA12         NP_06472           PDR15         NP_010694         CG11898         AAF56870         C47A10.1         CA803973         ABCA13         NP_689914           PDR5         NP_01773         CG14709         AAF56656         C54D1.1         AAC48149         ABCB7         AB005289           PXA2         NP_012713         CG1718         AAF50838         C54G10.3         CA481093         ABCC10         NP_258261           Sta6         NP_012713         CG1718         AAF50387         C5666.5         AAA81092         ABCC11         NP_01322           YCF1         NP_0103052         CG1810         AAF50366         F02E11.1         AA866050         ABCF1         AH34488           YDR061w         NP_010376         CG1819         AAF5047         F1286.1         AA864133         ABCF2         NP_00528           YEF8         P39376         CG1863         AAF50547         F1286.1         AA864133         ABCF3         ANP 04324           YER036C         NP_010336         CG2164         AAF5147         F1286.1         AA864133         ABCG3         AP20293           YER036C         NP_010376	PDRII	NP_012252	CG11460	AAF55727	C44B7.8	AAA68339	ABCA10	XP_085647
PDR15         NP_010694         CG11898         AAF56870         C47A10.1         CAB03973         ABCA13         NP_689914           PDR5         NP_014796         CG12703         AAF49018         CAB84.4         CA82384         ABCB5         AAO73470           PDR4         NP_012733         CG1470         AAF50828         C54G10.3         CAA99810         ABCB7         A0053297           PXA2         NP_012733         CG1703         AAF48069         C56E6.1         AAA81093         ABCC10         NP_258261           SNQ2         NP_010294         CG17338         AAF53736         DH11.3         CAA88940         ABCC12         NM_033226           SNQ5         CG17338         AAF53736         DH11.3         CAA88940         ABCC12         NP_03326           YCF1         NP_010376         CG1840         AAF50847         F1286.1         AA854153         ABCF2         NP_00583           YDR061w         NP_010376         CG1843         AAF59367         F186.4         CA393461         ABCG8         F320294           YER036C         NP_01182         CG1462         AAF5977         F2810.3         CAA82317         ABCA163         NP_01912           YR0307w         NP_01448         CG49247         <	PDR12	NP_015267	CG11897	AAF56869	C44B7.9	AAA68340	ABCA12	NP_056472
PDR5         NP_014796         CG12703         AAF49018         C4884.4         CAA8234         ABCB5         AAC7170           PXA1         NP_015178         CG14709         AAF54656         C5401.1         AAC48149         ABCB7         AB005289           PXA2         NP_010274         CG1703         AAF48069         C56E6.1         AAA81093         ABCC10         NP_14963           Ste6         NP_010713         CG1718         AAF50837         C56E6.5         AAA81092         ABCC11         NP_14963           YBT1         NP_010710         CG17646         AAF50837         C56E6.5         AAA81092         ABCC12         NP_14963           YDR6010         NP_010419         CG17646         AAF51341         E03G2.2         CAA88940         ABCC12         NP_03503           YDR091C         NP_010350         CG1819         AAF50847         F1264.1         AB66450         ABCF1         NP_005683           YER3         NP_010350         CG1813         AAF5360         F18E2.2         CAA93461         ABCG8         F320294           YFR030C         NP_010350         CG3164         AAF5367         F1986.4         CA8043217         ABCG 1         AC601945           YFR030V         NP_010950	PDR15	NP_010694	CG11898	AAF56870	C47A10.1	CAB03973	ABCA13	NP_689914
PXAI         NP_015178         CG14709         AAF5456         CS4D1.1         AAC48149         ABCB7         AB005289           PXA2         NP_012733         CG1494         AAF50838         CS4G10.3         CAA98103         ABCC10         NP_258261           SNQ2         NP_012713         CG1718         AAF50837         CS6E6.5         AAA81093         ABCC11         NP_149163           SYBC1         NP_0102713         CG1738         AAF5376         DH11.3         CAA89840         ABCC12         NM_033226           YCF1         NP_010419         CG16610         AAF50867         DF11.1         AA666050         ABCF1         AA143488           YDR061W         NP_010376         CG1814         AAF5047         F1286.1         AA854153         ABCF2         NP_060828           YEF3         NP_01350         CG182.4         AAF5164         F1986.4         CA89435         ABCG14         AF50293           YER03CC         NP_011282         CG186.3         AAF5148         F2086.3         AAA2317         ABCR (A4)         AF01945           YHR03CW         NP_011486         CG422.5         AAF5541         F22E10.1         CAA91800         BCR (Q1)         NP_01915           YNR070W         NP_01466	PDR5	NP_014796	CG12703	AAF49018	C48B4.4	CAA82384	ABCB5	AAO73470
PXA2NP_012733CGI494AAF50838CS4C103CAA88090ABCC10NP_021301SNQ2NP_012731CG1703AAF48069C56E6.1AAA81092ABCC10NP_128261Sre6NP_012731CG1738AAF50837C56E6.5AAA81092ABCC11NP_149163YBT1NP_013052CG17338AAF50376DH11.3CAA88940ABCC12NM_033226YCF1NP_010346CG1801AAF50836F02E1.1AAB64050ABCF1AH34488YDR091CNP_010376CG1819AAF50876F1286.1AAB54153ABCF2NP_006028yFF3NP_01330CG1824AAF49177F14F4.3CA854255ABCF3NP_060828yFR03CCNP_010953CG1824AAF50367F1986.4CAA93461ABCG8AF302934YHR03CNNP_011828CG1643AAF51548F2086.3AA82317ABCR (A4)AF001945Y1R03CNNP_011828CG3164AAF55767F22E10.2CAA91800BCR (G2)XP_032425Y0R01NNNP_014567CG452AAF5576F22E10.3CAA91801BSEP (G1)AF01952Y0R01NNNP_014567CG4561AAF5032F4221.1CAA91463MABC2 (B1)XP_032425Y0R01NNNP_014567CG4594AAF5323F5721.2CAA91801BSEP (G1)AF04591Y0R01NNNP_014567CG4594AAF5323F5721.2CAA91801BSEP (G1)AA56367Y0R01NNNP_014567CG4594AAF5323F5721.2 <td>PXAI</td> <td>NP_015178</td> <td>CG14709</td> <td>AAF54656</td> <td>C54D1.1</td> <td>AAC48149</td> <td>ABCB7</td> <td>AB005289</td>	PXAI	NP_015178	CG14709	AAF54656	C54D1.1	AAC48149	ABCB7	AB005289
SNQ2         NP_010294         CG1703         AAF48069         CS6E6.1         AAA81093         ABCC10         NP_258261           Ste6         NP_012713         CG1718         AAF50837         CS6E6.5         AAA81093         ABCC11         NP_149163           YBT1         NP_010419         CG17646         AAF51341         E03G2.2         CAA92148         ABCC13         NP_742021           YDR061w         NP_010376         CG1814         AAF50867         F02E11.1         AAB64050         ABCF2         NP_005683           YEF3         NP_010375         CG1824         AAF4177         F14F4.3         CAB54225         ABCG8         AF320294           YER03C         NP_010953         CG1824         AAF51548         F20B6.3         AAA82317         ABCG8         AF320294           YHR03C0         NP_010953         CG3164         AAF51548         F20B6.3         AAA82317         ABCR (A)         AF30294           YHR03V0         NP_011828         CG3164         AAF55767         F22E10.1         CAA91801         BSF (B11)         AF30294           YNR070w         NP_01468         CG4225         AAF55767         F22E10.3         CAA91801         BSF (B11)         AF91592           YOLN7C         NP_014	PXA2	NP_012733	CG1494	AAF50838	C54G10.3	CAA99810	ABCB9	AC002486
Ste6         NP_012713         CG1718         AAF50837         CS6E6.5         AAA81092         ABCC11         NP_149163           YBT1         NP_013052         CG17338         AAF53736         DH11.3         CAA892148         ABCC12         NM_033226           YCF1         NP_010419         CG17646         AAF50341         E03G2.2         CAA92148         ABCC13         NP_40201           YDR061w         NP_010376         CG1819         AAF50836         F02E11.1         AAB54153         ABCF2         NA4434488           YDR091C         NP_01330         CG1824         AAF6177         F14F4.3         CAA93451         ABCG8         AF32023           YER036C         NP_01933         CG18633         AAF53660         F18E2.2         CAA93451         ABCG8         AF32023           YER036C         NP_011828         CG164         AAF51548         F2086.3         AAA82317         ABCR (A4)         AF001945           YCR010W         NP_013030         CG327         AAF51726         F22E10.2         CAA91801         BCR (A1)         NP00195           YCN010w         NP_014687         CG4922         AAF55707         F22E10.2         CAA91801         BSEP (B1)         AF091582           YOR011w         NP_	SNQ2	NP_010294	CG1703	AAF48069	C56E6.1	AAA81093	ABCC10	NP_258261
YBT1         NP_013052         CG17338         AAF53736         DH11.3         CAA88940         ABCC12         NM_03226           YCF1         NP_010419         CG1746         AAF51341         E03G2.2         CAA8050         ABCC13         NP_742021           YDR061w         NP_010376         CG1801         AAF5047         F12Bc.1         AA86050         ABCF1         AH34488           YDR091C         NP_010376         CG1824         AAF5047         F12Bc.1         AA854153         ABCF3         NP_005683           YEF3         NP_013300         CG1824         AAF50360         F18E2.2         CA89421         ABCG8         AF30293           YER036C         NP_010933         CG3164         AAF5376         F1986.4         CA939461         ABCG8         AF30293           YHR0370w         NP_01828         CG3164         AAF5576         F12E1.0         CA802667         ALDP (D1)         CA79922           YNR070w         NP_01848         CG422.5         AAF55726         F22E10.1         CA91980         BCR (C2)         NP_01582           YOR011w         NP_878167         CG492.4         AAF5322         F22E10.3         CA91980         BCR (B0)         NP_0019515           YOR011w         NP_91797	Ste6	NP_012713	CG1718	AAF50837	C56E6.5	AAA81092	ABCCII	NP_149163
YCFI         NP_010419         CG17646         AAF51341         E03G2.2         CAA92148         ABCC13         NP_742021           YDR061w         NP_010376         CG1801         AAF50836         F02E11.1         AA866050         ABCF1         AAH34488           YDR091C         NP_010376         CG1819         AAF50847         F126.1         AA654135         ABCF2         NP_00583           yEF3         NP_010953         CG18633         AAF56360         F18E2.2         CAA99835         ABCG3         AF30294           YER03C         NP_0101830         CG3127         AAF5147         F196.4         CAA99341         ABCG8         AF30294           YHL03SC         NP_011828         CG3164         AAF51548         F206.3         AA82317         ABCR (A4)         AF01945           YKR103W         NP_014468         CG425         AAF5122         F216.4.2         CAA91800         BCR (A2)         NP_005155           YOL075C         NP_014567         CG4562         AAF55707         F22E10.1         CAA91800         BCR (A2)         NP_01582           YOR10W         NP_01457         CG4562         AAF5032         F22E10.3         CAA91800         CF1R (C7)         AAC16357           YDR1720W         NP_0	YBTI	NP_013052	CG17338	AAF53736	DHII.3	CAA88940	ABCC12	NM_033226
YDR061w         NP_010346         CG1801         AAF50836         F02E11.1         AAB66050         ABCF1         AAH3488           YDR091C         NP_010376         CG1819         AAF508047         F12B6.1         AAB54153         ABCF2         NP_005083           yEF3         NP_010350         CG1824         AAF6307         F14F4.3         CAB54225         ABC73         NP_006082           yEF8         P53978         CG18631         AAF53767         F1986.4         CAA93461         ABCG8         AF302294           YER036C         NP_010953         CG3164         AAF51548         F2086.3         AAA82317         ABCR (A4)         AF001945           YER030C         NP_011828         CG3164         AAF55767         F2210.1         CAA9160         BCRP (C2)         NP_001575           YCR070w         NP_014567         CG4522         AAF55726         F2210.1         CAA91800         BCRP (G2)         NP_032425           YOR011w         NP_07177         CG4822         AAF55326         F32210.1         CAA91802         CFTR (C7)         AAC13657           YOR011w         NP_011797         CG4822         AAF50312         F4210.1         AAA71121         MDR1 (B1)         4505769           YOR11w	YCFI	NP_010419	CG17646	AAF51341	E03G2.2	CAA92148	ABCC13	NP_742021
YDR091C         NP_010376         CG1819         AAF50847         F12B6.1         AAB54153         ABCF2         NP_005883           yEF3         NP_013350         CG1824         AAF48177         F14F4.3         CAB54225         ABCG3         NP_005883           yEF8         CG18033         AAF50360         F18E2.2         CAA99835         ABCG5         AF320294           YER036C         NP_010953         CG3164         AAF59367         F1986.4         CAA93461         ABCG2         AF302034           YER036C         NP_011828         CG3164         AAF59367         F1986.4         CAA91799         ALDP (D1)         CAA79222           YNR070w         NP_014567         CG4725         AAF55726         F22E10.1         CAA91799         ALDR (D2)         NP_005182           YOR011w         NP_014567         CG4794         AAF55726         F22E10.3         CAA91800         BCRP (G2)         XP_0032425           YOR1         NP_011797         CG4822         AAF50312         F42E11.1         CAA91802         CFTR (C7)         AAC13657           YOR1         NP_011797         CG4822         AAF50312         F42E11.1         CAA91803         MRE1 (B)         XP_001871           YPL226W         S65245	YDR061w	NP_010346	CG1801	AAF50836	F02E11.1	AAB66050	ABCFI	AAH34488
yEF3         NP_013350         CG1824         AAF48177         F14F4.3         CAB54225         ABCF3         NP_060828           yFF8         P53978         CG18633         AAF56360         F18E2.2         CAA99835         ABCG5         AF302291           YER036C         NP_010953         CG2316         AAF59367         F1986.4         CAA99836         ABCG8         AF302291           YHL035C         NP_011828         CG3164         AAF5122         F2164.2         CAB02667         ALDP (D1)         CAA799222           YNR070w         NP_014468         CG4225         AAF55241         F22E10.1         CAA91800         BCRP (G2)         XP_032425           YOR01w         NP_014567         CG4562         AAF55726         F22E10.2         CAA91800         BCRP (G2)         XP_035425           YOR1         NP_01797         CG4822         AAF55726         F22E10.4         CAA91802         CFTR (C7)         AAC13577           YP1226W         S65245         CG5651         AAF5322         F22E10.4         CAA91802         CFTR (C7)         AAC16357           YP1226W         S65245         CG5789         AAF5312         F42E1.1         CAA91802         MABC1 (B8)         AA607569           CG5584 <td< td=""><td>YDR091C</td><td>NP_010376</td><td>CG1819</td><td>AAF50847</td><td>F12B6.1</td><td>AAB54153</td><td>ABCF2</td><td>NP_005683</td></td<>	YDR091C	NP_010376	CG1819	AAF50847	F12B6.1	AAB54153	ABCF2	NP_005683
yEFB         P53978         CG18633         AAF56360         F18E2.2         CAA99835         ABCGS         AF320291           YER036C         NP_010953         CG2316         AAF59367         F1986.4         CAA93461         ABCG8         AF320294           YHL035C         NP_011828         CG3164         AAF51548         F2086.3         AAA82317         ABCG8         AF5021           YKR103W         NP_011828         CG3162         AAF51541         F2161.0         CAA91799         ALDP (D1)         CA47922           YNR070w         NP_014468         CG4225         AAF55241         F22E10.2         CAA91800         BCRP (G2)         XP_032425           YOR011w         NP_014567         CG452         AAF5532         F22E10.2         CAA91800         BCRP (G2)         XP_032425           YOR1         NP_011797         CG4822         AAF5332         F22E10.4         CAA91802         CFTR (C7)         AC13657           YPL226W         S65245         CG5653         AAF5332         F4221.1         CAA91430         MABC1 (B8)         AF07690           YPL226W         S65245         CG5789         AAF5332         F37C1.2         AAC71121         MDR1 (B1)         3505769           YPL226W         KG52	yEF3	NP_013350	CG1824	AAF48177	F14F4.3	CAB54225	ABCF3	NP_060828
YER036C         NP_010953         CG2316         AAF59367         F19B6.4         CAA93461         ABCG8         AF320294           YHL035C         NP_011828         CG3164         AAF51548         F20B6.3         AAA82317         ABCR (A4)         AF001945           YKR03W         NP_013030         CG3327         AAF51122         F21G4.2         CA802667         ALDP (D1)         CAA79922           YNR070w         NP_014468         CG4225         AAF55707         F22E10.1         CAA91900         BCP (G2)         XP_032425           YOL075C         NP_01467         CG4562         AAF55726         F22E10.3         CAA91801         BSEP (B1)         AF01552           YOR011W         NP_878167         CG4522         AAF55322         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YOR11W         NP_011797         CG4822         AAF5032         F42E11.1         CAA91463         MABC1 (B8)         AF047690           YPL226W         S65245         CG5581         AAF50312         F42E11.1         CAA91463         MABC1 (B8)         AAF0307           YPL226W         S65245         CG5651         AAF4305         F55G11.9         CAB0522         MDR1 (B1)         A36267           CG5658 <td>yEFB</td> <td>P53978</td> <td>CG18633</td> <td>AAF56360</td> <td>F18E2.2</td> <td>CAA99835</td> <td>ABCG5</td> <td>AF320293</td>	yEFB	P53978	CG18633	AAF56360	F18E2.2	CAA99835	ABCG5	AF320293
YHL035C         NP_011828         CG3164         AAF51548         F2086.3         AAA82317         ABCR (A4)         AF001945           YKR103W         NP_013030         CG3327         AAF51122         F21G4.2         CA802667         ALDP (D1)         CAA79922           YNR070w         NP_01468         CG4225         AAF5571         F22E10.1         CAA91799         ALDR (D2)         NP_005155           YOR070w         NP_014567         CG4562         AAF5576         F22E10.2         CAA91800         BSEP (B11)         AF03752           YOR011w         NP_878167         CG4794         AAF55726         F22E10.3         CAA91801         BSEP (B11)         AF047690           YOR1         NP_011797         CG4822         AAF5152         F22E10.4         CAA91802         CFTR (C7)         AAC14570           YPL226W         S65245         CG5513         AAF5032         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           YPL226W         S65245         CG5651         AAF5032         F55G11.9         CAB05222         MDR1 (B1)         4505769           CG5614         AAF5323         F57C12.4         AAB7072         MRP2 (C2)         CA65259           CG6162         AAF53328         K508F.9	YER036C	NP_010953	CG2316	AAF59367	F19B6.4	CAA93461	ABCG8	AF320294
YKR103W         NP_013030         CG3327         AAF51122         F21G4.2         CAB02667         ALDP (D1)         CAA79922           YNR070w         NP_014468         CG4225         AAF55241         F22E10.1         CAA91799         ALDR (D2)         NP_005155           YOL075C         NP_014567         CG4562         AAF5570         F22E10.2         CAA91800         BCRP (G2)         XP_032425           YOR011w         NP_07177         CG4922         AAF55726         F22E10.3         CAA91800         BSEP (B1)         AF03582           YOR1         NP_011797         CG4822         AAF5032         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5551         AAF50312         F42L1.1         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5581         AAF50312         F42L1.1         CAA91802         MBC1 (B8)         AF047690           CG5789         AAF5323         F35C11.9         CAB05222         MDR3 (B4)         AA36207           CG5051         AAF5323         F55G11.9         CAB0522         MDR3 (B4)         AA36207           CG6052         AAF5323         F57C12.5         AAB510355         MR91 (C1)<	YHL035C	NP_011828	CG3164	AAF51548	F20B6.3	AAA82317	ABCR (A4)	AF001945
YNR070w         NP_014468         CG4225         AAF55241         F22E10.1         CAA91799         ALDR (D2)         NP_005155           YOL075C         NP_014567         CG4562         AAF55707         F22E10.2         CAA91800         BCRP (G2)         XP_032425           YOR011w         NP_878167         CG4794         AAF55726         F22E10.3         CAA91801         BSEP (B1)         AF091582           YOR1         NP_011797         CG4822         AAF51552         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5651         AAF50342         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           CG5780         AAF52835         F43E2.4         AAC71121         MDR1 (B1)         A505769           CG5944         AAF49312         F56F4.6         AAB54203         MRP1 (C1)         AA846616           CG6162         AAF53233         F57C12.4         AAB07022         MR93 (G3)         AB010887           CG7346         AAF53328         K08E7.9         CAB01320         MRP4 (C4)         NP_005836           CG7491         AAF53268         T0201.5         CAB05909         MRP6 (C6)         AF076622           CG7491         AAF5	YKR103W	NP_013030	CG3327	AAF51122	F21G4.2	CAB02667	ALDP (D1)	CAA79922
YOL075C         NP_014567         CG4562         AAF55707         F22E10.2         CAA91800         BCRP (G2)         XP_032425           YOR011w         NP_878167         CG4794         AAF55726         F22E10.3         CAA91801         BSEP (B1)         AF091582           YOR1         NP_011797         CG4822         AAF51552         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5651         AAF50342         F42A10.1         AAA919072         MABC1 (B8)         AF047690           YPL226W         S65245         CG5651         AAF56312         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           CG5853         AAF52835         F43E2.4         AAC71121         MDR1 (B1)         4505769           CG5914         AAF5032         F55G1.6         CA80522         MDR3 (B4)         AA36207           CG6616         AAF5035         F57C12.4         AAB67022         MRP3 (C3)         AB010887           CG7346         AAF5035         F57C12.5         AAD31550         MRP4 (C4)         NP_005836           CG7491         AAF52639         T10H9.5         AAC19238         MTABC3 (B6)         NP_005680           CG7955         AAF4526	YNR070w	NP_014468	CG4225	AAF55241	F22E10.1	CAA91799	ALDR (D2)	NP_005155
YOR011w         NP_878167         CG4794         AAF55726         F22E10.3         CAA91801         BSEP (B11)         AF091582           YOR1         NP_011797         CG4822         AAF51552         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5651         AAF50342         F42A10.1         AAA19072         MABC1 (B8)         AF047690           YPL226W         S65245         CG5651         AAF50312         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           CG5789         AAF52335         F43E2.4         AAC71121         MDR1 (B1)         4505769           CG5944         AAF9305         F55G11.9         CA805222         MDR3 (B4)         AAA36207           CG6052         AAF49312         F56F4.6         AAB54203         MRP1 (C1)         AAB6616           CG6012         AAF53233         F57C12.4         AAB07022         MR93 (C3)         AB010887           CG7346         AAF50332         F57C12.5         AAD31550         MRP4 (C4)         NP_05836           CG7627         AAF53328         K08E7.9         CAB05929         MR96 (C6)         AF076422           CG7686         AAF552639         T10H.9.5         AAC19238	YOL075C	NP_014567	CG4562	AAF55707	F22E10.2	CAA91800	BCRP (G2)	XP_032425
YORI         NP_011797         CG4822         AAF51552         F22E10.4         CAA91802         CFTR (C7)         AAC13657           YPL226W         S65245         CG5651         AAF50322         F42A10.1         AAA19072         MABC1 (B8)         AF047690           CG5789         AAF56312         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           CG5853         AAF52835         F43E2.4         AAC71121         MDR1 (B1)         4505769           CG5052         AAF49305         F55G11.9         CAB05222         MDR3 (B4)         AAA36207           CG6052         AAF49312         F56F4.6         AAB5403         MRP1 (C1)         AAB46616           CG6162         AAF5035         F57C12.4         AAB07022         MR93 (C3)         AB010887           CG7346         AAF5035         F57C12.5         AAD31550         MRP4 (C4)         NP_005836           CG7627         AAF5328         K021.5         CAB05909         MR8C (G6)         AF076622           CG7806         AAF52639         T10H.5         AAC19238         MTABC3 (B6)         NP_005680           CG7795         AAF52639         T0H.5         AAC19218         MTABC3 (B6)         NP005746           CG7806	YOR011w	NP_878167	CG4794	AAF55726	F22E10.3	CAA91801	BSEP (BII)	AF091582
YPL226W         S65245         CG5651         AAF50342         F42A10.1         AAA19072         MABC1 (B8)         AF047690           CG5789         AAF56312         F42E11.1         CAA91463         MABC2 (B10)         XP_001871           CG5853         AAF52835         F43E2.4         AAC71121         MDR1 (B1)         4505769           CG5944         AAF49305         F55G11.9         CAB05222         MDR3 (B4)         AAA36207           CG6052         AAF49312         F56F4.6         AAB54203         MRP1 (C1)         AAB46616           CG612         AAF53223         F57C12.4         AAB07022         MR93 (C3)         AB010887           CG7346         AAF53223         F57C12.5         AAD31550         MRP4 (C4)         NP_005836           CG7491         AAF53328         K08E7.9         CAB01232         MR56 (C6)         AF076622           CG7627         AAF52639         T10H9.5         AAC19238         MTABC3 (B6)         NP_0058836           CG7955         AAF47525         T218.1         CAA94220         PMP6 (C6)         AF076422           CG8979         AAF58947         T2183         CAA94203         RNAse LI (E1)         CAS3972           CG89708         AAF57490         T26A5.1<	YORI	NP_011797	CG4822	AAF51552	F22E10.4	CAA91802	CFTR (C7)	AAC13657
CG5789       AAF56312       F42E11.1       CAA91463       MABC2 (B10)       XP_001871         CG5853       AAF52835       F43E2.4       AAC71121       MDR1 (B1)       4505769         CG5944       AAF49305       F55G11.9       CAB05222       MDR3 (B4)       AAA6207         CG6052       AAF49312       F56F4.6       AAB54203       MRP1 (C1)       AAB46616         CG6162       AAF56584       F57A10.3       CAB09418       MRP2 (C2)       CAA65259         CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF53328       K08E7.9       CAB01232       MRP4 (C4)       NP_005836         CG7491       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF58947       T21E8.2       CAA94203       RNAse L1 (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC7504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       C	YPL226W	S65245	CG5651	AAF50342	F42A10.1	AAA19072	MABCI (B8)	AF047690
CG5853       AAF52835       F43E2.4       AAC71121       MDR1 (B1)       4505769         CG5944       AAF49305       F55G11.9       CAB05222       MDR3 (B4)       AAA36207         CG6052       AAF49312       F56F4.6       AAB54203       MRP1 (C1)       AAB46616         CG6162       AAF56584       F57A10.3       CAB09418       MRP2 (C2)       CAA65259         CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF53328       K08E7.9       CAB01232       MRP4 (C4)       NP_005836         CG7491       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CA94220       PMP69 (D4)       AF009746         CG8473       AAF58947       T21E8.3       CAA94203       RNAse L1 (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AB802278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC6			CG5789	AAF56312	F42E11.1	CAA91463	MABC2 (B10)	XP_001871
CG5944       AAF49305       F55G11.9       CAB05222       MDR3 (B4)       AAA36207         CG6052       AAF49312       F56F4.6       AAB54203       MRP1 (C1)       AAB46616         CG6162       AAF56584       F57A10.3       CAB09418       MRP2 (C2)       CAA65259         CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF50355       F57C12.5       AAD31550       MRP4 (C4)       NP_005836         CG7491       AAF53328       K08E7.9       CAB01232       MRP6 (C6)       AAF07622         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94203       RNAse L1 (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC7504       SUR1 (C8)       AAB02278         CG9270       AAF33950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AA			CG5853	AAF52835	F43E2.4	AAC71121	MDRI (BI)	4505769
CG6052       AAF49312       F56F4.6       AAB54203       MRP1 (C1)       AAB46616         CG6162       AAF56584       F57A10.3       CAB09418       MRP2 (C2)       CAA65259         CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF50355       F57C12.5       AAD31550       MRP4 (C4)       NP_005836         CG7491       AAF53328       K08E7.9       CAB01232       MRP5 (C5)       AAB71758         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94203       PMP69 (D4)       AF009746         CG8473       AAF58947       T21E8.2       CAA94203       RNAse LI (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF4893       W04C9.1       AAC68724       TAP1 (B2)       CA440741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AA59841			CG5944	AAF49305	F55G11.9	CAB05222	MDR3 (B4)	AAA36207
CG6162       AAF56584       F57A10.3       CAB09418       MRP2 (C2)       CAA65259         CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF50035       F57C12.5       AAD31550       MRP4 (C4)       NP_005836         CG7491       AAF53328       K08E7.9       CAB01232       MRP5 (C5)       AAB71758         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF06622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF09746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA51416         CG8799       AAF53950       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CAA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AA59841			CG6052	AAF49312	F56F4.6	AAB54203	MRPI (CI)	AAB46616
CG6214       AAF53223       F57C12.4       AAB07022       MRP3 (C3)       AB010887         CG7346       AAF50035       F57C12.5       AAD31550       MRP4 (C4)       NP_005836         CG7491       AAF53328       K08E7.9       CAB01232       MRP5 (C5)       AAB71758         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA41416         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF485395       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AAA59841			CG6162	AAF56584	F57A10.3	CAB09418	MRP2 (C2)	CAA65259
CG7346       AAF50035       F57C12.5       AAD31550       MRP4 (C4)       NP_005836         CG7491       AAF53328       K08E7.9       CAB01232       MRP5 (C5)       AAB71758         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF09746         CG8473       AAF48511       T21E8.2       CAA94203       RNAse LI (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AAA59841			CG6214	AAF53223	F57C12.4	AAB07022	MRP3 (C3)	AB010887
CG7491       AAF53328       K08E7.9       CAB01232       MRP5 (C5)       AAB71758         CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA1416         CG8799       AAF58947       T21E8.3       CAA94203       RNAse LI (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF83950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AA59841			CG7346	AAF50035	F57C12.5	AAD31550	MRP4 (C4)	NP_005836
CG7627       AAF52648       T02D1.5       CAB05909       MRP6 (C6)       AF076622         CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA41416         CG8799       AAF58947       T21E8.3       CAA94203       RNAse LI (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AA59841			CG7491	AAF53328	K08E7.9	CAB01232	MRP5 (C5)	AAB71758
CG7806       AAF52639       T10H9.5       AAC19238       MTABC3 (B6)       NP_005680         CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA41416         CG8799       AAF58947       T21E8.3       CAA94203       RNAse LI (E1)       CA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CAA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AAA59841			CG7627	AAF52648	T02D1.5	CAB05909	MRP6 (C6)	AF076622
CG7955       AAF47525       T21E8.1       CAA94220       PMP69 (D4)       AF009746         CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA41416         CG8799       AAF58947       T21E8.3       CAA94203       RNAse LI (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CAA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AAA59841			CG7806	AAF52639	T10H9.5	AAC19238	MTABC3 (B6)	NP_005680
CG8473       AAF48511       T21E8.2       CAA94219       PMP70 (D3)       CAA41416         CG8799       AAF58947       T21E8.3       CAA94203       RNAse L1 (E1)       CAA53972         CG8908       AAF57490       T26A5.1       AAC77504       SUR1 (C8)       AAB02278         CG9270       AAF53950       T27E9.7       CAB04880       SUR2 (C9)       AF061323         CG9281       AAF48493       W04C9.1       AAC68724       TAP1 (B2)       CAA40741         CG9330       AAF49142       W09D6.6       CAB04947       TAP2 (B3)       AAA59841			CG7955	AAF47525	T21E8.1	CAA94220	PMP69 (D4)	AF009746
CG8799AAF58947T21E8.3CAA94203RNAse LI (E1)CAA53972CG8908AAF57490T26A5.1AAC77504SUR1 (C8)AAB02278CG9270AAF53950T27E9.7CAB04880SUR2 (C9)AF061323CG9281AAF48493W04C9.1AAC68724TAP1 (B2)CAA40741CG9330AAF49142W09D6.6CAB04947TAP2 (B3)AAA59841			CG8473	AAF485 I I	T21E8.2	CAA94219	PMP70 (D3)	CAA41416
CG8908AAF57490T26A5.IAAC77504SURI (C8)AAB02278CG9270AAF53950T27E9.7CAB04880SUR2 (C9)AF061323CG9281AAF48493W04C9.IAAC68724TAPI (B2)CAA40741CG9330AAF49142W09D6.6CAB04947TAP2 (B3)AAA59841			CG8799	AAF58947	T21E8.3	CAA94203	RNAse LI (EI)	CAA53972
CG9270AAF53950T27E9.7CAB04880SUR2 (C9)AF061323CG9281AAF48493W04C9.1AAC68724TAP1 (B2)CAA40741CG9330AAF49142W09D6.6CAB04947TAP2 (B3)AAA59841			CG8908	AAF57490	T26A5.1	AAC77504	SURI (C8)	AAB02278
CG9281         AAF48493         W04C9.1         AAC68724         TAPI (B2)         CAA40741           CG9330         AAF49142         W09D6.6         CAB04947         TAP2 (B3)         AAA59841			CG9270	AAF53950	T27E9.7	CAB04880	SUR2 (C9)	AF061323
CG9330 AAF49142 W09D6.6 CAB04947 TAP2 (B3) AAA59841			CG9281	AAF48493	W04C9.1	AAC68724	TAPI (B2)	CAA40741
			CG9330	AAF49142	W09D6.6	CAB04947	TAP2 (B3)	AAA59841

Alphabetic list, by taxon, of protein sequences used in this study								
C	G9663	AAF51130	Y39D8C.1	AAC69223	WHITE I (GI)	AAC51098		
C	G9664	AAF51131	Y39E4B.I	CAB54424	WHITE 2 (G4)	NP_071452		
C	G9892	AAF51223	Y42G9A.6	NP_498332				
C	G9990	AAF56807	Y43F8C.12	CAA21622				
М	dr49	AAF58437	Y47D3A.11	CAB57891				
М	dr50	AAF58271	Y48G8AL.11	AAK29911				
М	dr65	AAF50669	Y49E10.9	CAB11549				
Sc	arlet	AAF49455	Y50E8A.16	CAB60586				
Su	ır	AAF52866	Y53C10A.9	CAA22142				
N N	/hite	AAF45826	Y57GIIC.I	CAB16503				
			Y75B8A.26	CAA22110				
			ZK455.7	CAA91467				
			ZK484.2	AAK39394				

#### Table 2 (Continued)

recognizes substrate directly within the cytoplasmic leaflet of the plasma membrane [43], it does so at a much higher effective substrate concentration than would be the case if it recognized aqueous substrate. As a result, P-gp drug-binding sites can operate at relatively low affinity, and this, in turn, facilitates recognition of multiple substrates. This flexibility may be the key to explaining the range of tasks performed by ABC transporters, but also their apparently anomalous evolutionary history.

The mammalian P-gps include proteins capable of producing an MDR phenotype (MDR1 (B1)), as well as members with, apparently, specificity restricted to single physiological substrates such as phosphatidylcholine (MDR3 (B4)). As none of these have simple, orthologous, relationships with any of the C. elegans P-gps, no detailed predictions of function in nematode P-gps can be drawn on the basis of phylogeny alone. C. elegans P-gps do differ from one another in their ability to cause resistance to various environmental toxins [16], with no apparent correlation between phenotype and genetic distance from their mammalian homologs. Both human abca1 and nematode ced-7 mutants present similar apoptotic phenotypes, despite their rather distant relationship (Figure 3). ABCA1 mutations also cause defects in high-density lipoprotein cholesterol transport, and it is still an open question as to whether the analogous function of these two homologs in apoptosis accurately predicts a sharing of other functions. Similar limitations on the extent to which function may be predicted from sequence alone are likely to obtain in those subfamilies whose members are noted for variability and multiplicity of function, that is, subfamilies A, B, C and G.

Schriml and Dean [10] speculated that the distinct clustering of amino- and carboxy-terminal halves of ABCA proteins suggests that full ABC transporters have generally evolved from half-transporters. The pattern of structural change within the closely related subfamilies ABCD, ABCC and ABCB does suggest that the half-transporter configuration was the ancestral one for at least these three subfamilies (Figure 2). It also reveals instances where half-transporters have evolved from duplicated genes, as in the origination of ABCB from a fragment of an ABCC gene, and that, in turn, some ABCB genes have duplicated again, in giving rise to the P-gp genes.

A comprehensive comparison of worm and yeast genomes [23] noted that while most of the nematode genome did not closely resemble that of yeast, there was a strongly conserved 20% of the nematode genome that had a high degree of homology to a corresponding 40% of the yeast genome. Within this highly conserved subset of genes, there was a very frequent finding of orthology between members of the two genomes. As many as 57% of the most closely related gene pairs contained exactly one worm and one yeast gene. The obvious inference is that one corresponding gene was present in the common ancestor of the two species. Their overall picture of genome evolution is one in which a conserved cadre of proteins performs core biological functions required by all eukaryotes. These would remain essentially invariant throughout eukaryotes, and one expects analogous functions to be carried out by orthologous genes across large evolutionary distances. These gene families are presumably protected over the long run by their essential and irreplaceable roles in basic biochemical functions required by all organisms. However, as Chervitz et al. [23] point out, only a minority of gene families fit this mode, with most genes belonging to poorly conserved or taxonomically restricted families.

We expected that the frequency of simple orthologous gene pairs typical of highly conserved gene families shared by both yeast and worm would hold true for our comparison between nematode and human versions of such a highly conserved gene family as ABC transporters. However, this generality Table 3

Frequency of orthologous pairs among ABC transporters							
	Sc	Ce	Dm	Hs			
Sc		10%	17%	10%			
Ce	3		14%	16%			
Dm	5	8		22%			
Hs	5	8	П				

Numbers below the diagonal represent the number of orthologous pairs of ABC transporters, according to our phylogeny, found in pairwise comparisons between each of the four genomes in this study. Percentages above the diagonal are calculated from the corresponding number given for that pair, divided by the smaller of the two counts of ABC transporters in that pair of genomes. Ce, *C. elegans*; Dm, *D. melanogaster*; Hs, *H. sapiens*; Sc, *S. cerevisiae*.

clearly does not apply to ABC transporters, despite their strong conservation across all domains of life. It seems reasonable to suppose that the rather loose relationship between substrate specificity and amino acid sequence that characterizes ABC transporters allows for much more potential exchange and sorting of biological functions among homologous genes than is typical. In turn, this pervasive preadaptation for functional overlap enables organisms to survive the occasional loss of substantial numbers of ABC transporters and to rapidly re-evolve lost functionality by coopting homologous genes.

The evolutionary dynamic we propose here is reminiscent of an explanation put forward by Huynen et al. [44] to explain a pattern observed in a comparative analysis of 11 microbial genomes. They found that the frequency distribution of genefamily sizes within each completely sequenced genome tended to follow a power-law distribution across a 30-fold range of genome sizes. Their model is one in which genes are duplicated or deleted randomly in time, but the gene families are coherent with respect to the probability of duplication or deletion in each time unit in the simulation. In other words, the probability of duplicating or deleting a gene may change over time, but every member of a gene family always has the same probability of duplication or deletion as every other member of the family. So, whereas a given family can be either favored for expansion or targeted for deletion in a given time period, all members of the family are equally favored or disfavored by selection at the same time. Huynen et al. argued that this property of 'dynamic coherence' in a gene family could arise if all gene-family members have more or less the same function, so that they are all favored or disfavored by selection at the same time, depending on how much that function is needed.

Under a power-law distribution, gene families would tend to be subject to fluctuations of a size on the same order as the gene-family size itself [44]. We should then expect that

typical gene families will have undergone very substantial episodes of expansion and near-extinction, and in Huynen et al.'s model all gene families do become extinct within a finite time. It is evident that ABC transporters are highly atypical for a strongly conserved gene family, in that the family as a whole is highly conserved across genomes despite being subject to the same large fluctuations in size, which would tend to eventually eliminate gene families whose members are not individually indispensable. It should be noted that the ABC family does not seem uniformly subject to one or the other mode of evolution. Subfamilies E and F, which are not involved with transport, but rather have roles in translation and gene regulation, fit the 'strongly conserved' [23] model very well, retaining simple orthologous relationships over long spans of time. Only the transporter subfamilies themselves, because of their highly adaptable substrate-recognition capability, are subject to large fluctuations in size. We propose that finding large sets of paralogous genes, and infrequently conserved orthologs, in a gene family reflects ongoing cycles of gene loss and reacquisition of analogous functions in distantly related, newly expanded, lineages. Furthermore, we suggest that this is in fact the expected outcome of dynamic coherence, a mode shared, perhaps, by most of the less-conservative gene families, as well as the ABC genes.

We expect that future functional studies, to determine the extent of parallel and convergent evolution among ABC transporters, will eventually allow us to discern the fundamental roles of ABC transporters that ensure their long-term survival as a group. Also of interest will be whether the functional suites of genes fulfilling these roles are bounded in any way that resembles the phylogenetic subdivisions into which we presently categorize these proteins.

## Materials and methods Identification of ABC transporter genes

A computer file, WormPep16 [45], containing 16,332 protein sequences predicted from the completed C. elegans genome was searched using the FASTA program [46]. Our initial query sequences were those of known C. elegans ABC proteins (for example, Pgp-1, the D. melanogaster white gene homolog T26A5.1, and so on). Matching protein sequences returned by FASTA were checked by BLAST [47], using either the NCBI [48] or Baylor College of Medicine (BCM) servers [49]. Only those with highly significant matches to annotated ABC proteins in the sequence database were retained. The most poorly matched, verified ABC protein from each FASTA run was used as the query sequence for an additional FASTA search, and this process was repeated until no new ABC proteins were found. At a later stage in the analysis, representative members of different ABC transporter subfamilies were used as query sequences to search the updated WormPep81 file using a BLAST server at the Sanger Centre [45]. Searches were conducted using multiple queries until all proteins already included in our dataset were found. No additional ABC proteins were identified, though some sequences were found to have been included in our dataset twice under different names. These redundant sequences were eliminated. FASTA searches were run on a SUN Microsystems UltraSPARC 5 computer. All other computer operations were carried out on an Apple Power Macintosh G3. Yeast and human ABC transporter sequences were obtained from NCBI and are described in the literature [10,13].

#### Identification of ABC protein features

BLAST + Beauty searches on the BCM server identified the location of the conserved Walker A and ABC signature motifs (Prosite motifs [50] PS00017 and PS00211, respectively) associated with the ATP-binding cassette(s) of each protein. The number and positions of transmembrane domains in each ABC protein were predicted by using TopPred II v1.3 [51] and then vetting the program's results by eye to exclude spurious transmembrane segments. Chromosomal locations of each ABC protein in the *C. elegans* genome were looked up in the *C. elegans* database AceDB [52].

#### Phylogenetic analyses

Using the information derived from each protein sequence (as above) we extracted only the sequence of each predicted ATP-binding cytoplasmic domain. These domains were assembled into a single file using the SeqApp1.9 multiple sequence editor [53], and aligned using ClustalX [54]. In those cases where two ATP-binding cassettes (ABCs) are present in a single protein with no intervening transmembrane domains (Subfamilies E and F, see Figure 1), the entire sequence was divided into two at an arbitrary point halfway between the two predicted ABC domains. As a result, 'twodomain' proteins are represented twice in our initial analysis. Once this approach had been used to assign genes to particular well-supported subgroups, we realigned the sequences and reanalyzed the relationships within each group using fulllength amino acid sequence data.

Aligned sequences were used to generate matrices of mean distances between proteins, and these matrices were used to generate phylogenetic trees according to the neighbourjoining algorithm [55], refined using the SPR branch-swapping technique under the minimum evolution criterion, implemented by PAUP\*4.0b10 [56]. Bootstrapping (1,000 replicates) was done according to the method of Felsenstein [57], using the same parameters described above. Phylogenetic trees were visualized and manipulated using TreeView 1.6.2 [58] and MacClade 3.0.4 [59].

#### Additional data files

The following additional data are included with the online version of this article: the protein sequence alignments for the ABCA subfamily (Additional data file 1), the ABCB subfamily (Additional data file 2), the ABCC subfamily (Additional data file 3), the ABCD subfamily (Additional data file 4), the ABCE and ABCF subfamilies (Additional data file 5), the ABCG subfamily (Additional data file 6), the ABCH subfamily (Additional data file 7), and the protein sequences from the nucleotide-binding folds only (Additional data file 8). In addition to the four genomes discussed in this paper, mouse (*M. musculus*) ABC transporter genes are included in some of these alignments. All eight files are in Nexus format, which is a plain-text format designed for use with the programs PAUP [56] and MacClade [59]. A Nexus Data Editor for Windows is also available [60].

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