Functional Analogy in Human Metabolism: Enzymes with Different Biological Roles or Functional Redundancy?

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Accepted: July 4, 2017

Abstract

Since enzymes catalyze almost all chemical reactions that occur in living organisms, it is crucial that genes encoding such activities are correctly identified and functionally characterized. Several studies suggest that the fraction of enzymatic activities in which multiple events of independent origin have taken place during evolution is substantial. However, this topic is still poorly explored, and a comprehensive investigation of the occurrence, distribution, and implications of these events has not been done so far. Fundamental questions, such as how analogous enzymes originate, why so many events of independent origin have apparently occurred during evolution, and what are the reasons for the coexistence in the same organism of distinct enzymatic forms catalyzing the same reaction, remain unanswered. Also, several isofunctional enzymes are still not recognized as nonhomologous, even with substantial evidence indicating different evolutionary histories. In this work, we begin to investigate the biological significance of the cooccurrence of nonhomologous isofunctional enzymes in human metabolism, characterizing functional analogous enzymes identified in metabolic pathways annotated in the human genome. Our hypothesis is that the coexistence of multiple enzymatic forms might not be interpreted as functional redundancy. Instead, these enzymatic forms may be implicated in distinct (and probably relevant) biological roles.

Key words: enzymatic activity, convergent evolution, H. sapiens.

De Novo Origin of Enzymatic Activities and Functional Analogy in Human Metabolism

Enzymes have their biological activities defined by the type of chemical transformation carried out and by the mechanism through which this reaction is executed. The chemical transformations accomplished by enzymes are classified using the recommendations of the Nomenclature Committee of the International Union of Biochemistry (http://www.chem.qmul.ac.uk/iubmb/enzyme/). Based on the reaction catalyzed by the enzyme an Enzyme Commission (EC) number is assigned. According to this hierarchical classification, each enzyme receives a 4-digit number: the first digit describes the general chemical reaction catalyzed by the enzyme (the enzyme class); the two subsequent numbers have different meanings depending on the class of the enzyme; the fourth digit describes the specificity of the reaction, defining the specific substrate/product or cofactors used (McDonald and Tipton 2014).

In silico comparisons of metabolic pathways predicted from completely sequenced genomes of a variety of prokaryotic and eukaryotic species revealed incomplete or even absent pathways in several organisms (Cordwell 1999; Galperin and Koonin 1999; Huynen et al. 1999; Morett et al. 2003; Peregriń-Alvarez et al. 2003; Hanson et al. 2010). In some of these cases, the “missing” enzymes were replaced by functional equivalent molecules, able to catalyze the same reaction but exhibiting virtually no similarity in their amino acid chains, thus escaping identification by methods based on sequence similarity. These nonhomologous isofunctional molecules, known as analogous enzymes, arise from independent evolutionary events, converging for the same biological function, and may be associated with both related or unrelated phylogenetic lineages and/or possess different catalytic mechanisms, as well as distinct fold topologies and three-dimensional (3D) structures (Cordwell 1999; Galperin and Koonin 1999; Huynen et al. 1999; Morett et al. 2003; George et al. 2004; Gherardini et al. 2007; Omelchenko et al. 2010).

Several studies have suggested that the fraction of enzymatic activities in which multiple events of independent origin have taken place during evolution is substantial (Hegyi and
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Gerstein 1999; Morett et al. 2003; George et al. 2004; Gherardini et al. 2007; Omelchenko et al. 2010), and some of these “missing” enzymes have been identified and characterized in some detail (Almonacid et al. 2010; Galperin and Koonin 2012). Apparently, analogous enzymes are often recruited from distinct superfamilies (Galperin et al. 1998; Omelchenko et al. 2010), with some of these alternative forms sharing the reaction catalyzed and the configuration of the catalytic residues (although these residues do not share the same fold, in these cases) (Galperin and Koonin 2012).

Despite being recognized for a long time, though erroneously referred in older literature as isozymes (or isoenzymes), isoforms, or class/type I and class/type II enzymes (e.g., Martin and Schnarrenberger 1997), functionally analogous enzymes remain poorly explored, and a comprehensive investigation of the occurrence, distribution, and implications of convergence in enzymatic activities, at least involving organisms whose genomes have been completely sequenced, has not been done so far. Fundamental questions, such as how analogous enzymes originate, why so many events of independent origin have apparently occurred during evolution, and what are the reasons for the coexistence in the same organism of distinct enzymatic forms catalyzing the same biochemical reaction, among several other questions, such as concerning the catalysis of similar reactions by different structural scaffolds (Almonacid et al. 2010), remain unanswered.

Surprisingly, numerous isofunctional enzymes are still not recognized as nonhomologous counterparts, despite substantial evidence indicating different evolutionary histories (e.g., Omelchenko et al. 2010). However, in some of these unrecognized cases, it has been demonstrated that the analogous enzymes either have an unsuspected separate evolutionary history or present (experimentally verified) distinct functional features, as we will discuss later.

In this work, we begin to investigate the biological significance of the cooccurrence of nonhomologous isofunctional enzymes in human metabolism, characterizing functional analogous enzymes identified in biochemical pathways and processes annotated in the human genome. Our hypothesis is that the coexistence of multiple enzymatic forms might not be interpreted as functional redundancy. Instead, these enzymatic forms may be implicated in distinct (and probably relevant) biological roles.

To catalog the repertoire of isofunctional enzymes cooccurring in the human metabolism (from now on referred as intragenomic analogous enzymes) a computational pipeline (AnEnPi) (Otto et al. 2008) was employed to identify putative analogous enzymes using the KEGG database (Kanehisa and Goto 2000) as the source of information. The predicted functional analogy instances were confirmed based on domain, folding and 3D structure information assigned to the enzymes implicated (see Materials and Methods for details).

Altogether, we could find evidence of convergence in 15 enzymatic activities belonging to 45 distinct processes and metabolic pathways represented in KEGG’s human reference maps (table 1, and supplementary materials I and II, Supplementary Material online). The genomic coordinates of the genes encoding these predicted analogous enzymes showed that these genes are dispersed throughout the human genome, with most of the genes encoding for distinct analogous forms (as well as duplications of several alternative forms) located on separated chromosomes (fig. 2).

The Repertoire of Nonhomologous Isofunctional Enzymes in Humans

One valuable source of information on enzymatic activities and metabolic pathways is the KEGG Pathway database available at the Kyoto Encyclopedia of Genes and Genomes (KEGG) platform, which comprises a collection of manually elaborated maps representing the current knowledge about networks of molecular interaction in biological processes or biochemical pathways. Hence, from KEGG database version 73.1, we obtained 1,159,633 protein sequences encoded in 2,494 genomes, ranging all three domains of life (Archaea, Bacteria, and Eukarya), distributed in 3,825 enzymatic activities. From these enzymatic activities, 3,572 were fully annotated with the four EC digits classification, containing 1,025,885 protein sequences. On the other hand, 253 incomplete ECs were identified (defined until the first, second, or third digit of the EC classification scheme), comprising 133,748 sequences.

Different types of convergence occur at the molecular level and can be categorized into functional, mechanistic, structural, and sequence. Thus, enzymes whose chemical transformations are defined only by three digits of EC classification may have different reaction specificities (different substrates/products or cofactors), constituting mechanistic analogous, that is, unrelated enzymes which catalyze distinct chemical transformations through the same mechanism of action (Doolittle 1994; Gherardini et al. 2007). However, this sort of event is not considered in this work, which is dedicated solely to investigate functional analogy. Thus, the AnEnPi computational prediction (including all organisms and enzymatic activities in KEGG database) resulted in 2,203 enzymatic activities in which protein sequences were grouped in two or more distinct clusters, comprising 1,996 enzymatic activities with four-digit EC annotation.

Considering only the inference of convergence in enzymatic activities with four-digit EC classification annotated in the human genome, we found 150 ECs (2,288 protein sequences) identified by AnEnPi as sustaining putative events of de novo origin. After removing from our data set enzymatic activities in which protein sequences were annotated as “subunits” and “chains,” as well as enzymatic activities containing clusters composed of a single human sequence or in
## Table 1

Sequence and Structure Similarity Profile of Protein Sequences Comprising Each Enzymatic Activity Assigned to the Bona Fide (+) Data Set of Intragenomic Analogous Enzymes

| EC       | Gene UniprotKB | PDB Cluster | Gene UniprotKB | PDB Cluster | Identity (%) | Similarity (%) | Score | TM-Score | RMSD |
|----------|----------------|-------------|----------------|-------------|--------------|----------------|-------|----------|------|
| 1.3.1.20 | DHDH Q9UQ10    | 2O48        | AKR1C2 P52895  | 2HDJ        | 16.3         | 28.0           | 40.0  | 0.34186  | 6.09 |
|          | AHX Q9UQ10     | 2O48        | AKR1C1 Q04828  | 1J96        | 16.0         | 27.5           | 33.0  | 0.33888  | 5.89 |
| 1.15.1.1 | SOD2 P04179    | 1LVV        | SOD1 P00441    | 4XCR        | 13.6         | 24.8           | 37.5  | 0.24203  | 4.48 |
|          | SOD2 P04179    | 1LVV        | SOD3 P08294    | 2JLP        | 3.7          | 6.7            | 21.5  | 0.28352  | 5.09 |
| 2.4.1.22 | B4GALT2 O60909 | 1LUV        | B4GALT1 P15291 | 2AH9        | 50.0         | 62.7           | 104.95| ND       | ND   |
| 1.15.1.1 | SOD2 P04179    | 1LVV        | B4GALT1 P15291 | 2AH9        | 15.4         | 23.7           | 30.5  | ND       | ND   |
| 2.4.2.31 | SIRT6 Q8N6T7   | 3K35        | SIRT6 Q8N6T7   | 3K35        | 15.4         | 23.7           | 30.5  | ND       | ND   |
| 2.7.1.67 | PI4KA P42356   | 3T9A        | PI4K2A Q9BTU6  | 4XCR        | 10.8         | 17.1           | 52.7  | ND       | ND   |
| 2.7.4.21 | PIP5K2 Q6PFW1  | 3T9A        | PIP5K2 Q6PFW1  | 3T9A        | 56.2         | 64.5           | 417.05| ND       | ND   |
| 3.1.1.3  | AADAC P22760   | 3CEL        | LIPC P11150    | 3CEL        | 10.3         | 17.7           | 100.5 | ND       | ND   |

(continued)
| EC | Gene | UniprotKB | PDB | Cluster | Gene | UniprotKB | PDB | Cluster | Identity (%) | Similarity (%) | Score  | TM-Score | RMSD |
|----|------|---------|-----|--------|------|---------|-----|--------|------------|-------------|--------|----------|------|
|    | LIPC | P11150  | ND  | 4      | PNLI| P54315  | 2PPL| 4      | 29.0       | 43.9        | 506.0  | ND       | ND    |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 41.3       | 61.0        | 1059.5 | ND       | ND    |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 47.3       | 63.7        | 1113.5 | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| Q17RR3  | ND  | 4      | 30.5       | 42.2        | 556.0  | ND       | ND    |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 64.0       | 79.7        | 1676.0 | 0.94537  | 1.37  |
|    | PNLI| P54315  | 2PPL| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
|    | PNLI| P16233  | 1LPB| 4      | PNLI| P54315  | 2PPL| 4      | 67.3       | 80.6        | 1750.0 | 0.93392  | 1.76  |
|    | LIPC | P11150  | ND  | 4      | PNLI| P11150  | ND  | 4      | 27.1       | 43.0        | 473.5  | ND       | ND    |
Table 1 Continued

| EC   | Gene UniprotKB | PDB  | Cluster Gene UniprotKB | PDB  | Cluster | Identity (%) | Similarity (%) | Score | TM-Score | RMSD |
|------|----------------|------|------------------------|------|---------|--------------|----------------|-------|----------|------|
| 3.1.3.5 | NT5C1B Q96P26 ND | 2 | NT5C1A Q9BXI3 ND | 2 | 35.4 | 43.0 | 114.0 | ND | ND |
| NT5C1B Q96P26 ND | 2 | NTSE P21589 4H2G | 3 | 8.5 | 13.3 | 26.0 | ND | ND |
| NT5C1A Q9BXI3 ND | 2 | NTSE P21589 4H2G | 3 | 10.8 | 20.1 | 32.5 | ND | ND |
| NT5C1B Q96P26 ND | 2 | NTSC Q8TCD5 4LS7 | 5 | 7.3 | 13.2 | 29.5 | ND | ND |
| NT5C1A Q9BXI3 ND | 2 | NTSC Q8TCD5 4LS7 | 5 | 9.7 | 15.1 | 29.0 | ND | ND |
| NT5C1B Q96P26 ND | 2 | NTSCA Q9H0P0 2CN1 | 7 | 8.6 | 16.6 | 35.5 | ND | ND |
| NT5C1A Q9BXI3 ND | 2 | NTSCA Q9H0P0 2CN1 | 7 | 9.1 | 17.1 | 23.5 | ND | ND |
| 3.1.4.12 | SMPD2 O60906 ND | 1 | SMPD3 Q9NY59 ND | 1 | 10.4 | 15.2 | 87.5 | ND | ND |
| SMPD2 O60906 ND | 1 | SMPD1 P17405 5I81 | 2 | 5.5 | 10.0 | 35.5 | ND | ND |
| SMPD3 Q9NY59 ND | 1 | SMPD1 P17405 5I81 | 2 | 2.6 | 3.6 | 51.0 | ND | ND |
| SMPD2 O60906 ND | 1 | SMPD4 Q9NXE4 ND | 3 | 8.2 | 13.6 | 62.0 | ND | ND |
| SMPD3 Q9NY59 ND | 1 | SMPD4 Q9NXE4 ND | 3 | 12.7 | 20.2 | 51.5 | ND | ND |
| SMPD2 O60906 ND | 1 | ENPP7 Q6UWV6 5UDIO | 4 | 7.0 | 12.8 | 27.5 | ND | ND |
| SMPD3 Q9NY59 ND | 1 | ENPP7 Q6UWV6 5UDIO | 4 | 13.3 | 19.2 | 29.5 | ND | ND |
| SMPD1 P17405 5I81 | 2 | SMPD4 Q9NXE4 ND | 3 | 6.3 | 10.7 | 39.0 | ND | ND |
| SMPD1 P17405 5I81 | 2 | ENPP7 Q6UWV6 5UDIO | 4 | 16.1 | 25.0 | 82.5 | 0.2898 | 6.61 |
| SMPD4 Q9NXE4 ND | 3 | ENPP7 Q6UWV6 5UDIO | 4 | 5.6 | 8.7 | 49.5 | ND | ND |
| 4.2.99.18 | NTHL1 P78549 ND | 1 | OG11 O15527 1KO9 | 1 | 19.0 | 28.4 | 85.5 | ND | ND |
| NTHL1 P78549 ND | 1 | NEIL2 Q96F92 1VZP | 1 | 14.6 | 25.7 | 29.0 | ND | ND |
| NTHL1 P78549 ND | 1 | NEIL1 Q96F92 1TDH | 1 | 6.7 | 9.8 | 40.0 | ND | ND |
| OG11 O15527 1KO9 | 1 | NEIL2 Q96F92 1VZP | 1 | 2.9 | 4.1 | 21.0 | 0.2789 | 4.84 |
| OG11 O15527 1KO9 | 1 | NEIL1 Q96F92 1TDH | 1 | 14.3 | 20.2 | 42.5 | 0.28166 | 6.65 |
| NTHL1 P78549 ND | 1 | APEX1 P72695 2O3H | 1 | 15.6 | 23.0 | 41.0 | ND | ND |
| NTHL1 P78549 ND | 1 | APEX2 Q9U8Z4 ND | 1 | 4.7 | 6.6 | 14.5 | ND | ND |
| OG11 O15527 1KO9 | 1 | APEX1 P72695 2O3H | 1 | 16.5 | 26.7 | 23.5 | 0.27998 | 6.74 |
| OG11 O15527 1KO9 | 1 | APEX2 Q9U8Z4 ND | 1 | 9.4 | 14.6 | 42.5 | ND | ND |
| NTHL1 P78549 ND | 1 | APLF Q81W19 2KUO | 6 | 3.7 | 6.1 | 24.5 | ND | ND |
| OG11 O15527 1KO9 | 1 | APLF Q81W19 2KUO | 6 | 6.0 | 8.7 | 20.5 | 0.14656 | 6.94 |
| NEIL2 Q96F52 1VZP | 1 | NEIL1 Q96F41 1TDH | 1 | 18.8 | 24.9 | 143.5 | 0.50213 | 2.44 |
| NEIL2 Q96F52 1VZP | 1 | APEX1 P72695 2O3H | 1 | 13.2 | 22.2 | 19.5 | 0.28098 | 4.80 |
| NEIL2 Q96F52 1VZP | 1 | APEX2 Q9U8Z4 ND | 1 | 6.6 | 10.5 | 42.0 | ND | ND |
| NEIL1 Q96F41 1TDH | 1 | APEX1 P72695 2O3H | 1 | 5.2 | 7.3 | 44.0 | 0.25323 | 6.80 |
| NEIL1 Q96F41 1TDH | 1 | APEX2 Q9U8Z4 ND | 1 | 14.6 | 20.8 | 41.0 | ND | ND |
| NEIL2 Q96F52 1VZP | 1 | APLF Q81W19 2KUO | 6 | 3.9 | 7.9 | 24.5 | 0.09438 | 5.32 |
| NEIL1 Q96F41 1TDH | 1 | APLF Q81W19 2KUO | 6 | 10.7 | 18.3 | 46.5 | 0.15631 | 6.56 |
| APEX1 P72695 2O3H | 1 | APEX2 Q9U8Z4 ND | 1 | 14.9 | 21.0 | 264.0 | ND | ND |
| APEX1 P72695 2O3H | 1 | APLF Q81W19 2KUO | 6 | 6.7 | 12.8 | 47.5 | 0.16022 | 6.62 |
| APEX2 Q9U8Z4 ND | 1 | APLF Q81W19 2KUO | 6 | 12.3 | 20.5 | 44.5 | ND | ND |
| 5.3.99.2 | PTGDS P41222 2WWP | 1 | HPGD5 O60760 1YI | 2 | 13.7 | 21.4 | 9.5 | 0.28983 | 5.76 |
| 5.3.99.3 | PTGES2 Q9H772 ND | 1 | PTGES O14684 4AL0 | 2 | 0.6 | 0.6 | 18.0 | ND | ND |

*The 3D model of the human dehydrogenase (UniProt Q9UQ10) was obtained using the crystal structure of a Macaca fascicularis dehydrogenase (PDB 2048) by comparative modeling.*
which the protein sequences were grouped in one single cluster (see Materials and Methods for details), we obtain 116 protein sequences comprising 32 distinct enzymatic activities in human metabolism.

A flowchart representing our downstream data analyses is shown in figure 1. Overall, 116 protein sequences were initially predicted as pairs or groups of alternative forms in 32 enzymatic activities of the human metabolism. From these, 70 protein sequences, comprising 15 ECs, were assigned to the bona fide (+) data set, in which all enzymatic activities are composed of putative alternative enzymatic forms belonging to at least two distinct superfamilies. The remaining 46 sequences (17 ECs), assigned to the (−) data set, were rejected from our analysis.

Hydrolase class (36 protein sequences in 6 ECs) was the most frequent class in our bona fide (+) data set, followed by Transferases (17 protein sequences in four ECs), Oxidoreductases (six protein sequences in two ECs), Lyases (seven protein sequences in one EC), and Isomerases (four protein sequences in two ECs). No evidence of convergence was found in Ligase class. On the other hand, these 15 enzymatic activities are mapped in 45 biochemical pathways or processes of several major metabolic classes: Aging, Cancers, Carbohydrate metabolism, Cellular community—eukaryotes, Development, Digestive system, Endocrine system, Glycan biosynthesis and metabolism, Immune diseases, Lipid metabolism, Metabolism of cofactors and vitamins, Metabolism of other amino acids, Neurodegenerative diseases, Nucleotide metabolism, Replication and repair, Signal transduction, Transport and catabolism, Xenobiotics biodegradation and metabolism (supplementary material I, Supplementary Material online).

It is worth noticing that 12 of these 15 enzymatic activities in the bona fide (+) data set (~73%) were previously reported as presenting evidence of analogy (Capriles et al. 2010; Omelchenko et al. 2010): 1.3.1.20 (Trans-1,2-dihydrobenzene-1,2-diol dehydrogenase), 1.15.1.1 (Superoxide dismutase), 2.7.4.21 (Inositol-hexakisphosphate kinase), 3.1.1.29 (Aminoacyl-IRNA hydrolase), 3.1.2.2 (Palmitoyl-CoA hydrolase), 3.1.3.2 (Acid phosphatase), 3.1.3.5 (5’-nucleotidase), 3.1.4.12 (Sphingomyelin phosphodiesterase), 4.2.99.18 (DNA-(apurinic or apyrimidinic site) lyase), 5.3.99.2 (Prostaglandin-D synthase), 5.3.99.3 (Prostaglandin-E synthase), and 3.1.1.3 (Triacylglycerol lipase).

The SUPERFAMILY database (Wilson et al. 2009) consists of a collection of hidden Markov models, representing structural protein domains according to SCOP superfamily classification. Consequently, a superfamily groups together domains which...
have an evolutionary relationship. Hence, considering the SUPERFAMILY classification, we identified 39 different superfamilies (38 distinct folds) among the putative analogous enzymes in the bona fide (+) data set. The most frequent superfamilies are: alpha/beta-Hydrolases (13), Phosphoglycerate mutase-like (6), Lipase/lipoxygenase domain (PLAT/LH2 domain) (6), ADP-ribosylation (5), DNase I-like (4), HAD-like (4), Metallo-dependent phosphatases (3), SAICAR synthase-like (3), NAD(P)-linked oxidoreductase (2), Protein kinase-like (PK-like) (2), Cu,Zn superoxide dismutase-like (2), DNA-glycosylase (2), Glutathione synthetase ATP-binding domain-like (2), GST C-terminal domain-like (2), Nucleotide-diphospho-sugar transferases (2), S13-like H2TH domain (2), and Thioredoxin-like (2), followed by 22 different superfamilies represented once. On the other hand, we identified 51 distinct Pfam domains/families in those 70 enzymes. Of these, 29 enzymes are multidomain and 41 are composed of (or annotated as) a single domain. Three domains are shared among some enzymatic activities: His_Phos_2 (ECs 2.7.4.21 and 3.1.3.2), Metallophos (ECs 3.1.3.2, 3.1.3.5 and 3.1.4.12), and Exo_endo_phos (ECs 3.1.4.12 and 4.2.99.18). With two exceptions, enzymatic forms assigned to separate AnEnPi clusters have correspondingly different domain composition, indicating that inside a particular cluster-EC group, sequences might share a common origin.

However, alternative forms of the enzymatic activity 2.7.1.67 display the same domain composition (PI3_PI4_kinase), although they are members of unrelated superfamilies (ARM repeat, Protein kinase-like (PK-like), and ADP-ribosylation). Enzymatic activity 4.2.99.18, on the other hand, exhibits a much more complex pattern of domain and superfamily composition (supplementary material I, Supplementary Material online).

To measure the similarity among these 70 sequences in the bona fide (+) data set, we performed a global rigorous pairwise sequence alignment (table 1). The highest score, similarity, and identity values were observed between enzymatic forms belonging to the same AnEnPi cluster, as expected, since enzymes that share the same enzymatic activity, grouped in the same cluster, are presumably homologous.

We obtained similar results when 3D structures of these protein sequences were compared, employing the TM-score (Zhang and Skolnick 2004) and RMSD (root-mean-square deviation of atomic positions) measurements to estimate the similarity between them. We applied the following thresholds to distinguish related and unrelated structures: TM-score < 0.2, indicating a probable distinct evolutionary origin, and TM-score > 0.5, mostly corresponding to the same fold in SCOP (Murzin et al. 1995) or CATH (Sillitoe et al. 2015). Most of the alternative forms obtained TM-scores < 0.5.
when their structures were aligned (table 1 and supplementary material II, Supplementary Material online). Therefore, comparisons between sequences belonging to the same AnEnPi cluster-EC group resulted in RMSD values tending to zero and TM-scores close to 1, indicating a possible common evolutionary origin. When sequences belonging to distinct clusters of the same EC were compared, the opposite trend was observed, as expected (table 1). The intermediate TM-scores observed between the products of the genes NT5C3A and the alternative forms encoded by genes NT5C (0.50279) and NT5M (0.50929), as well as between the products of the genes CEL and LIPF (0.47684), can be attributed to the folds HAD-like and alpha/beta-Hydrolases shared between them, respectively.

In summary, we could assess the inference of convergence in all those 15 enzymatic activities based on superfamily and domain information, and based on structural alignments between the predicted alternative forms in 9 out of 15 of those enzymatic activities as well (ECs 1.3.1.20, 1.15.1.1, 2.4.1.22, 2.7.1.67, 3.1.2.2, 3.1.3.2, 3.1.3.5, 4.2.99.18, and 5.3.99.2). As shown in figure 2A, except for genes PTGES and PTGES2 encoding enzymes of the enzymatic activity 5.3.99.3, on chromosome 9 the genes encoding intragenomic analogous enzymes appear to be randomly distributed, dispersed throughout the entire human genome and recognized in 21 of the 24 nuclear chromosomes (20 autosomes and one sex chromosome). For genes encoding alternative forms as well as genes encoding homologous enzymatic forms, we mapped the chromosomal locations and then plotted in a circular diagram, as shown in figure 2B. Likewise, the distances between genes encoding intragenomic analogous and between homologous enzymes, exhibit a similar fuzzy pattern of occurrence in the human genome.

**Nucleotidases, Dehydrogenases, Synthases, Dismutases, Kinases, and Lipases**

The literature indicates the existence of seven human 5'-nucleotidases (EC 3.1.3.5), hydrolases involved in the biosynthesis of nucleosides and inorganic phosphate from noncyclic nucleoside monophosphates, encoded by the genes NT5E, NT5C1A, NT5C1B, NT5C, NT5C3A, NT5C2, and NT5M: one soluble enzyme associated with the cell membrane (NT5E), and six enzymes with an intracellular location, either cytosolic (NT5C1A, NT5C1B, NT5C, NT5C3A, and NT5C2) or mitochondrial (NT5M) (Zukowska et al. 2015). All these genes are distributed in several distinct chromosomes (1, 2, 6, 7, 10, and 17) (fig. 2), and the enzymes encoded by them were assigned to five AnEnPi clusters: 1) NT5E, 2) NT5C1A and NT5C1B, 3) NT5C and NT5M, 4) NT5C3A, and 5) NT5C2 (supplementary material I, Supplementary Material online).

The enzyme encoded by the gene NT5E belongs to the 5'-nucleotidase, C-terminal domain, and Metallo-dependent phosphatases superfamily, whereas the one encoded by the gene NT5C, as well as most of the remaining human 5'-nucleotidases (NT5C3A, NT5C2, NT5M), are members of the HAD-like superfamily. The enzymes encoded by the genes NT5C1A and NT5C1B do not have any superfamily annotation or any available 3D structure but were grouped together in a separate AnEnPi cluster showing considerable sequence similarity, indicating a possible common origin (table 1). The membrane-bound enzyme, NT5E, clearly distinguishes from the remaining enzymatic forms in all measures, as it was allocated in a separate AnEnPi cluster, showing remarkably low sequence and structural similarity when compared with all other 5'-nucleotidases, an entirely different superfamily/fold classification (as mentioned before), and a distinct domain composition/architecture (table 1 and supplementary material I, Supplementary Material online). On the other hand, the cytosolic, HAD-like superfamily enzymes, encoded by genes NT5C, NT5C3A, and NT5C2, as well as the mitochondrial enzyme, encoded by the NT5M gene, were assigned to three separate AnEnPi clusters; NT5C and NT5M enzymes, residing in the same AnEnPi cluster, show high sequence and structural similarity between them, as well as the same domain composition/architecture, whereas the opposite trend is observed when HAD-like superfamily enzymes representatives of distinct AnEnPi clusters are compared (both NT5C or NT5M against NT5C3A or NT5C2, and NT5C3A against NT5C2): very low sequence and structural similarity, and unrelated domain composition/architecture (table 1 and supplementary material I, Supplementary Material online).

Interestingly, Crisp et al. (2015) showed considerable evidence that genes NT5C and NT5M had been horizontally acquired in the human lineage (possibly from bacterial genomes), therefore contributing to biochemical diversification of 5'-nucleotidases during animal evolution. Besides the diversity of subcellular localization, possible evolutionary origin, amino acid sequence, fold, and domain composition/architecture, these enzymes use 5'-nucleotides from various sources, displaying significant differences in the range of substrates (partially overlapping), as well as in substrate specificity (Zimmermann 1992). Hence, it is reasonable to think of the possibility of these enzymes fulfill different biological roles while regulating diverse physiological processes.

The oxidoreductases Trans-1,2-Dihydrobenzene-1,2-Diol Dehydrogenase (EC 1.3.1.20) comprises the enzymes encoded by the genes DHDH, AKR1C1, and AKR1C2. In our analyses, AnEnPi assigned the product of the gene DHDH to a separate cluster, whereas all remaining enzymes (encoded by AKR1C1 and AKR1C2 genes) were grouped in a different cluster. The predicted alternative forms could be distinguished based on domain composition/architecture, superfamily classification, as well as 3D structure (table 1, and supplementary material I and II, Supplementary Material online). It is worth noticing that DHDH gene is located on chromosome 19, whereas the remaining genes are all neighbors,
colocated in chromosome 10, with their products presenting almost identical amino acid sequences (97.8% identity over the entire sequences), therefore reinforcing the evidence of common origin (possibly recent duplication) for the genes AKR1C1 and AKR1C2 (fig. 2). The low sequence and structural similarity between DHDH enzyme and members of the aldo-keto reductase family (e.g., AKR1C1 and AKR1C2 enzymes) has already been reported, as well as differences in use of substrates (Arimitsu et al. 1999; Carbone et al. 2008). DHDH enzyme acts on (–)[1R,2R]-dihydrodiols, while aldo-keto reductases oxidize (+)[15,25]-dihydrodiols (Carbone et al. 2008). Also, aldo-keto reductase members use synthetic steroids as substrate (Penning et al. 2015). We are aware that homologous enzymes can also present distinct substrate specificities, but in this case, the established substrate difference clearly correlates with the assumed separate evolutionary origin, even in the absence of further information that could indicate other possible implication(s) in distinct biological roles.

Representatives of the enzymatic class isomerase, prostaglandin D2 synthase and hematopoietic prostaglandin D synthase (encoded by the genes PTGDS, located on chromosome 9, and HPGDS, located on chromosome 4, respectively) (EC 5.3.99.2), both regulate the synthesis of prostaglandin D2, acting in signaling and inflammatory processes (Trimarco et al. 2014; Urade and Eguchi 2002; Lim et al. 2013). Our computational pipeline AnEnPi assigned PTGDS and HPGDS enzymes to separate clusters, and subsequent analyses revealed that these enzymes are also unrelated based on domain composition, superfamily classification, as well as amino acid sequence and 3D structure (table 1, and supplementary materials I and II, Supplementary Material online), we confirmed that these enzymes are indeed unrelated, corroborating previous evidence of functional convergence in SOD enzymatic activity (Omelchenko et al. 2010). In a recent study, Garcia et al. (2017) demonstrated that manganese-dependent enzymes with superoxide dismutase activity, SodA and SodM, not only coexist in the human pathogen Staphylococcus aureus but also clearly display distinct biological roles, in which solely one of the alternative forms, SodM, can promote resistance to antibiotics and host immunity. The authors showed that SodA is strictly manganese-dependent and relevant for combating oxidative stress as well as for disease development when manganese is abundant, whereas SodM is truly cambialistic, essential under manganese-deplete conditions, maintaining equal enzymatic activity in the presence of manganese or iron (Garcia et al. 2017). Even though this phenomenon has only been demonstrated in bacteria so far, it opens the opportunity to explore it in other prokaryotic or eukaryotic species.

Members of the class transferase, enzymes with 1-phosphatidylinositol 4-kinase activity (PI4Ks) (EC 2.7.1.67) participate in inositol phosphate metabolism and phosphatidylinositol signaling system, catalyzing the phosphorylation of phosphatidylinositol. The product of this reaction is phosphatidylinositol 4-phosphate, a primary precursor in the synthesis of phosphatidylinositolpolyphosphates, molecules involved in many biologic processes, such as signal transduction, membrane trafficking, and cytoskeletal reorganization (Barylko et al. 2001). The mammalian PI4Ks have been classified into two types, II and III, based on biochemical characteristics, and the literature highlights the existence of different domain organizations between PI4Ks of type II (genes PI4K2A and PI4K2B) and PI4Ks of type III (genes PI4KA and PI4KB), with PI4KA and PI4KB being more similar to each other, and PI4KA bearing a characteristic binding domain (Boura and Nencka 2015; Heilmeyer et al. 2003). Hence, the division of human PI4Ks in two separate AnEnPi clusters of putative isofunctional nonhomologous forms—one of these clusters formed by the products of the genes PI4KA and PI4KB, and the other one comprising PI4K2A and PI4K2B gene products, corresponding to the mammalian PI4Ks of type II and III, respectively, as well as their assignment to distinct superfamilies (except for the enzyme encoded by the gene PI4K2B which has no superfamily classification) and unrelated 3D structures (supplementary materials I and II,
Supplementary Material online), reinforces similar results obtained in previous studies concerning this enzymatic activity (reviewed by Boura and Nencka 2015).

Overall, the all-against-all pairwise sequence comparison among protein sequences of each enzymatic activity of the bona fide (+) data set corroborated the AnEnPi computational predictions. However, we found at least two cases in which AnEnPi’s clustering method may have “produced” more human enzymatic forms than expected: EC 3.1.3.5 (S’-nucleotidase), with five clusters, and EC 3.1.1.3 (Triacylglycerol Lipase), with four clusters. In the EC 3.1.3.5, enzymes encoded by the genes NT5C, NT5M, NT5C3A, and NT5C2 (cytoplasmic forms) share the same superfamily class (HAD-like), whereas the enzyme encoded by the gene NT5E (membrane form) is simultaneously classified in two superfamilies: S’-nucleotidase (syn. UDP-sugar hydrolase), C-terminal domain and Metallo-dependent phosphatases. Similarly, the products of the genes AADAC, CEL, LIPC, PNLIP, PNLIPRP1, PNLIPRP3, LIPG, and LIPF, comprising the EC 3.1.1.3, are all assigned to the alpha/beta-Hydrolases superfamily, whereas the product of the PNPLA3 gene belongs to a distinct superfamily (FabD/lyso phospholipase-like). Another piece of evidence supporting this assumption is that the genes PNLIPRP1, PNLIPRP2, PNLIPRP3, PNLIP, and LIPF are all neighbors, located on human chromosome 10, and their corresponding enzymes share considerably higher sequence similarity among them, than with PNPLA3, possibly representing duplication events (fig. 2 and table 1).

As we expected, all nonhomologous iso-functional enzymes we could characterize are assigned to distinct KEGG orthologous groups (KO; http://www.genome.jp/kegg/ko.html), corroborating the distinct evolutionary origin for the predicted alternative forms (supplementary material I, Supplementary Material online). Only six KOs are shared between two or more sequences in our bona fide (+) data set: K01081, grouping six out of seven S’-nucleotidases (NT5C1B, NT5C1A, NT5C, NT5M, NT5C3A, and NT5C2); K01046, including two out of ten triacylglycerol lipases (LIPC, and LIPG); K01068, gathering three out of five palmitoyl-CoA hydrolases (ACOT2, ACOT1, and ACOT4); K07756 and K13024 containing all five inositol-hexakisphosphate kinases (IP6K1, IP6K3, IP6K2, and PPIPK2, PPIPK1, respectively); and K13711, grouping two out of four 1-phosphatidylinositol 4-kinases (PI4K2A and PI4K2B).

After an extensive literature search, we were unable to find further information that could indicate (or not) a distinct evolutionary origin for the alternative forms of the remaining nine enzymatic activities of our bona fide (+) data set, neither their possible implication in distinct biological roles: EC 2.4.1.22 (Lactose synthase), EC 2.4.2.31 (NAD+—protein-arginine ADP-ribosyltransferase), EC 2.7.4.21 (Inositol-hexakisphosphate kinase), EC 3.1.1.29 (Aminoacl-TRNA hydrolase), EC 3.1.1.13 (Triacylglycerol lipase), EC 3.1.2.2 (Palmitoyl-CoA hydrolase), EC 3.1.3.2 (Acid phosphatase), EC 3.1.4.12 (Sphingomyelin phosphodiesterase), EC 4.2.99.18 (DNA- (apurinic or apyrimidinic site) lyase), and EC 5.3.99.3 (Prostaglandin-E synthase).

Different Biological Roles or Functional Redundancy?

In this work, we found substantial evidence of nonhomologous iso-functional enzymes coexisting in 15 enzymatic activities (comprising 70 enzymatic sequences) of human metabolism. Notably, despite the use of very restrictive criteria (excluding multimeric enzymes, enzymatic activities with incomplete EC classification, as well as clusters composed exclusively of a single human sequence) and our focus on human enzymatic activities in which the participation of unrelated enzymes are recognized, we discovered intragenomic analogous enzymes in three enzymatic activities (20% of our bona fide data set) with no evidence of analogy reported so far: lactose synthase (EC 2.4.1.22), NAD+—protein-arginine ADP-ribosyltransferase (EC 2.4.2.31), and 1-phosphatidylinositol 4-kinase (EC 2.7.1.67). These enzymatic activities participate in nine distinct biochemical pathways or biological processes, some of which playing essential roles in cancer, galactose metabolism, glycosaminoglycan biosynthesis, glycoprophilipid biosynthesis, inositol phosphate metabolism, mannose type O-glycan biosynthesis, n-glycan biosynthesis, other types of o-glycan biosynthesis, and phosphorylidyinositol signaling system.

We hypothesize that the coexistence of multiple nonhomologous iso-functional enzymes in the human metabolism might not be interpreted as functional redundancy since these intragenomic analogous enzymes might be implicated in distinct biological roles. To test this hypothesis, we will be comparing the transcription profile of the genes encoding the repertoire of intragenomic analogous enzymes cataloged in human metabolism, using RNA-Seq data obtained from 8,555 samples of 53 distinct healthy human tissues publicly available at the GTEx portal (The GTEx Consortium 2013)(http://www.gtexportal.org/home/). The identification of alternative enzymatic forms differentially expressed or coexpressed could provide evidence regarding possible distinct biological roles played by human intragenomic analogous enzymes.

Materials and Methods

Computational Prediction of Analogy

Protein sequences from 2,494 completely sequenced genomes comprising organisms of the three domains of life were obtained from the KEGG database release 73.1 (Kanehisa and Goto 2000) and clustered by enzymatic activity, based on the degree of similarity between their amino acid sequences, applying the methodology described in Otto et al. (2008), implemented in AnEnPi pipeline; sequences sharing the same
enzymatic activity but assigned to two or more distinct clusters are considered putative functional analogous, indicating one or more possible events of independent evolutionary origin.

Briefly, we compared 1,159,633 enzymatic sequences, separately by enzymatic activity (EC), all against all, using BLAST+ version 2.2.30 (Altschul 1997) and default parameters. Next, we transformed the sequence alignment result in a graph in which each enzymatic sequence represents a node. For each enzymatic activity, any sequence (node) pair that achieved an alignment score \( \geq 120 \) were connected by an edge; linked sequences are presumably homologous, therefore were grouped in the same AnEnPi cluster; on the other hand, enzymatic sequences grouped in distinct AnEnPi clusters of the same enzymatic activity are presumably analogous. The number of subgraphs obtained represents the number of putative events of independent origin in each enzymatic activity or, in other words, the number of times a particular enzymatic activity has arisen during evolution. The similarity threshold used in the clustering phase (BLAST score \( \geq 120 \)) is based on a significant experimental observation: enzymes that proved to share the same enzymatic activity and present significantly different 3D structures (based on structural alignments), scored below 120 when their amino acid sequences were compared with BLAST (Galperin et al. 1998). Although the absence of detectable sequence similarity might often be attributed to the divergence between homologous sequences during evolution, it was observed that many alternative forms of enzymes catalyzing the same biochemical reaction had significantly distinct 3D structures, and therefore have (presumably) evolved independently (Galperin et al. 1998; Omelchenko et al. 2010).

Subsequently, the AnEnPi output was processed as follows: 1) incomplete ECs were removed. Enzymes whose chemical transformations are defined only up to the third digit of the EC classification may have different reaction specificities (different substrates/products or cofactors) and, in this case, the predicted analogues would correspond to a mechanistic analogy. However, this type of analogy is not part of the scope of this work, which is devoted exclusively to the study of functional analogy; 2) enzymatic activities in which enzymes were annotated as “subunit” and “chain” were manually inspected and excluded, because the presence of heteromultimeric enzymes in the data set can inflate the number of analogy events detected. This problem arises during the process of annotation of enzymatic sequences, in which different subunits (or chains) of a multimeric enzyme often inherit the annotated activity for the enzyme as a whole disregarding its evolutionary origin and its participation in the related activity; 3) enzymatic activities containing clusters composed exclusively of a single human sequence were removed. If a single sequence distinguishes from tens or hundreds of other enzymatic sequences (from humans and/or other species) we consider it suspicious, as this might represent functional misannotation; 4) enzymatic activities in which the occurrence of alternative enzymatic forms was not detected (composed of a single cluster) were removed.

Validation of Predicted Intragenomic Analogy in Human

We used protein domains, superfamily/folding, and 3D structure annotations retrieved for proteins within and between clusters of each enzymatic activity to confirm putative cases of analogy detected inside the human genome (intragenomic analogy). These data were obtained from Pfam 27.0 (Finn et al. 2014) (http://pfam.xfam.org/) and SUPERFAMILY 1.75 (Wilson et al. 2009) (http://supfam.org/SUPERFAMILY).

Experimentally resolved 3D structures for proteins were retrieved from PDB database (Berman et al. 2000) (http://www.rcsb.org/) and previous information of convergence in enzymatic activities were selected from the scientific literature (Omelchenko et al. 2010). Based on superfamily classification, we split our data set into two data sets: \((+)\) and \((-)\) data set. Enzymatic activities composed of putative alternative enzymatic forms belonging to at least two distinct superfamilies were assigned to the \((+)\) data set, otherwise were assigned to the \((-)\) data set.

For the bona fide \((+)\) data set, we performed an all-against-all pairwise sequence comparison among all sequences inside each enzymatic activity using the optimal global sequence alignment implemented in the software Needle (Rice et al. 2000).

We generated structural models for sequences without 3D information employing the comparative modeling software Modeller (Webb and Sali 2014). For this, we used templates from PDB database retrieved by BLAST similarity searches (coverage in query > 70%; coverage in subject > 90%; identity > 30%; e-value < 10-3). Modeller generated 50 structural models and the best model for each protein was selected based on the lowest DOPE (Discrete Optimized Protein Energy) score value. Subsequently, the quality of these selected models was evaluated with SAVES (http://services.mbi.ucla.edu/SAVES/) and MolProbity (Chen et al. 2010). The side chains were fitted with KiNG (Chen et al. 2009), and the energy minimization was performed with ModRefiner (Xu and Zhang 2011).

The 3D structures were generated with PyMOL (The PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC) and the protein structural alignments were performed with TM-align (Zhang and Skolnick 2005). RMSD values and TM-scores (Zhang and Skolnick 2004) were calculated with the TM-align package. TM-score distances were normalized by the average size of the chains of each compared structure.

Genomic and Metabolic Mapping of Analogous Enzymes

Genomic coordinates of genes encoding the alternative forms in the bona fide \((+)\) data set were retrieved from Ensembl
(genome version: GRCh38) (Cunningham et al. 2015). The ideogram representing the chromosomal localization of these genes was created with the software PhenoGram (http://ritchielab.psu.edu/). Additionally, a circular diagram displaying the genomic distances between genes encoding alternative forms (distinct AnEnPi cluster of the same EC) as well as genes encoding homologous enzymatic forms (belonging to the same AnEnPi cluster and EC) was created with Circos (Krzywinski et al. 2009).

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

**Acknowledgments**

We wish to thank Fernando Alvarez-Valin, Marcelo Alves Ferreira, and Leandro Mattos Pereira for fruitful discussions; André Luiz Quintanilha Torres and Luiz Philippe Ribeiro Baptista for their help in the analysis. We also thank to Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) and the Plataforma de Bioinformática da Fiocruz RPT04A/RJ for their support.

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Associate editor: Bill F. Martin