# HbVar Database of Human Hemoglobin Variants and Thalassemia Mutations: 2007 Update

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HbVar (http://globin.bx.psu.edu/hbvar) is a locus-specific database (LSDB) developed in 2001 by a multi-center academic effort to provide timely information on the genomic sequence changes leading to hemoglobin variants and all types of thalassemia and hemoglobinopathies. Database records include extensive phenotypic descriptions, biochemical and hematological effects, associated pathology, and ethnic occurrence, accompanied by mutation frequencies and references. In addition to the regular updates to entries, we report significant advances and updates, which can be useful not only for HbVar users but also for other LSDB development and curation in general. The query page provides more functionality but in a simpler, more user-friendly format and known single nucleotide polymorphisms in the human  $\alpha$ - and  $\beta$ -globin loci are provided automatically. Populationspecific β-thalassemia mutation frequencies for 31 population groups have been added and/or modified and the previously reported  $\delta$ - and  $\alpha$ -thalassemia mutation frequency data from 10 population groups have also been incorporated. In addition, an independent flat-file database, named XPRbase (http://www.goldenhelix.org/xprbase), has been developed and linked to the main HbVar web page to provide a succinct listing of 51 experimental protocols available for globin gene mutation screening. These updates significantly augment the database profile and quality of information provided, which should increase the already high impact of the HbVar database, while its combination with the UCSC powerful genome browser and the ITHANET web portal paves the way for drawing connections of clinical importance, that is from genome to function to phenotype. © 2007 Wiley-Liss, Inc.

KEY WORDS: LSDB; globin genes; thalassemia; variants; mutation screening; software

### INTRODUCTION

Hemoglobinopathies are the most common inherited disorders in humans, resulting from mutations in the  $\alpha$ and  $\beta$ -globin gene clusters (reviewed in Forget et al., 2001). The  $\alpha$ -globin gene cluster is composed of the genes
HBZ (MIM# 142310), HBA2 (MIM# 141850), HBA1 (MIM# 141800), HBM (MIM# 609639) and HBQ1 (MIM#

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142240), which encode the  $\zeta$ -,  $\alpha$ 2-,  $\alpha$ 1-, and possibly  $\mu$ - and  $\theta$ -globin polypeptides, respectively. The  $\beta$ -globin gene cluster is composed of the genes *HBE1* (MIM# 142100), *HBG2* (MIM# 142250), *HBG1* (MIM# 142200), *HBG1* (MIM# 14200), *HBG1* (MIM# 14200), which encode the  $\epsilon$ -,  ${}^{G}\gamma$ -,  ${}^{A}\gamma$ -,  $\delta$ - and  $\beta$ -globin polypeptides, respectively. Single nucleotide substitutions can lead to hemoglobin (Hb) variants, due to amino acid replacements, while molecular defects in either regulatory or coding regions of the human  $\alpha$ -,  $\beta$ - or  $\delta$ -globin genes can minimally or drastically reduce their expression, leading to  $\alpha$ -,  $\beta$ - or  $\delta$ - thalassemia respectively.

In 2001, we developed HbVar, derived from previous compilations (Huisman et al., 1997, 1998), as a publicly available locus-specific database (LSDB) to provide timely information to interested users, e.g. the globin research community, providers of genetic services and counseling, patients and their parents, pharmaceutical industries, *etc.* New hemoglobin variants and thalassemias continue to be discovered, and thus HbVar was designed for regular entry updates and corrections. The query interface provides easy access to this information for the research community and for physicians as an aid in diagnosis. HbVar has rapidly become an important aid in the globin research community and is considered to be one of the premier LSDBs available to date (Claustres et al., 2002, Beroud, 2005). The major advantages of HbVar compared to other LSDBs, are the mutation information quality and depth of coverage, its regular updates and upgrades, and, most importantly, its interrelation with other databases, such as GALA (Giardine et al., 2003), GenPhen and PhenCode (Giardine et al., submitted). Minor disadvantages were the lack of information on the available experimental protocols to screen for globin gene mutations and the database's lengthy query page.

We report here several new updates in HbVar's structure and contents, aiming at increasing the quality of the database and its impact to society. Also, the possible implications of these updates in the curation of other LSDBs are discussed.

#### UPDATES TO EXISTING DATA

HbVar's information has been expanded by more than 400 additional entries and corrections, made continually by the database curators during the two recent upgrades (Table 1). In order to identify new Hb variants and thalassemia mutations not previously documented in the database, HbVar records were compared against Online Mendelian Inheritance in Man (MIM) records and external links were placed in each HbVar entry, redirecting the user to the respective MIM entry. Also, articles from the specialized journal *Hemoglobin*, which frequently publishes new Hb variants and thalassemia mutations, were manually scanned and where applicable, previously undocumented mutations have been included into HbVar. Although empirical nomenclature is widely used in HbVar, proper [Human Genome Variation Society (HGVS); http://www.hgvs.org] mutation nomenclature is also automatically provided next to each entry, based on the reference sequence.

As far as population-specific data are concerned,  $\beta$ -thalassemia mutation frequencies for 31 population groups have been added and/or modified where needed, while the previously reported  $\delta$ - and  $\alpha$ -thalassemia mutation frequency data from 10 population groups have also been extracted from the published literature and made available for the HbVar users..

Finally, we have recently introduced a unique identifier for each HbVar entry. This feature will be particularly useful for curators to avoid duplicate entries and also for users when referring to a particular mutant, especially those lying within mutational hotspots.

#### QUERY PAGE UPDATES AND NEW FUNCTIONALITIES

The previous query page for HbVar was lengthy, so in order to streamline it without losing any functionality, we have grouped all previous options by type and included them in 13 different hyperlinked sections (Fig. 1).

# HbVar: A database of Human Hemoglobin Variants and Thalassemias

Query Page		
Name:		
	Type of Thalassemia	I:
Category: Any		beta0 beta+
Chain: ☐ Agamma ☐ Gga	mma □ alpha1 □ alpha2 □ beta □ delta □ zeta1 □ zeta2	_
	R □ exon □ intron □ not within known transcription unit	
Montallan data	•	
Mutation data	a □ No mutation data in database	
wissing mucacion data	No mutation data in database	Range
Deed her words on	Choose a numbering system	
Residue number:	Common DNA-based description	From
Substitutions Insertions	e reference sequence from an image	То
<u>Deletions</u> Fusion gene Hbs		
Contact		
Haplotype		
Hematology		
Electrophoresis		
<u>Chromatography</u>		
Stability		
Ethnic background	_	
Aborigine		
Acadian Afghan		
African	▼	
Frequency range from	to %	
Group tested: Size from	to , Description	
Occurrence comments		
Structure studies		
Functional studies		
Comments Onery for SNPs at the by	man globin loci using UCSC Browsers	
Query for Sixes at the nu	man groom foci using ocise browsers	
Alpha globin region, UC		
References		
Within boxes combined with Ol Submit Query	R, between boxes with AND	

**Figure 1.** The new HbVar query page. All of the features of the previously lengthy query page (Hardison et al., 2002) have been incorporated in the various hyperlinks. Also, new functionalities have been added for screening for SNPs and for graphically displaying the reported mutants per region for every gene (see text for details).

Clear form

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Therefore, if a user wishes to query the database based on a variant's occurrence, he/she only needs to open the "Occurrence" section and narrow the search by selecting the relevant criteria included therein. This minimize/maximize option has improved the clarity of display of the query page. Also, a "query history" has been added, which displays a list of the queries a user has previously run. Tools on this page not only enable the user to retrieve previous query results but also to perform operations on the results, such as unions, intersections and subtractions, to refine them.

**Table 1.** Summaries of Mutation Categories in the HbVar Database for Hemoglobin Variants and Thalassemia Mutations (Note That Some Entries Appear in More Than One Category)

Date	Count of results			
Entries	October 2001 <sup>a</sup>	October 2006 b		
Entries involving the <i>HBA1</i> gene (OMIM# 141800)	212	254		
Entries involving the <i>HBA2</i> gene (OMIM# 141850)	250	295		
Entries involving the <i>HBB</i> gene (OMIM# 141900)	635	711		
Entries involving the <i>HBD</i> gene (OMIM# 142000)	57	72		
Entries involving the <i>HBG2</i> gene (OMIM# 142200)	44	49		
Entries involving the <i>HBG1</i> gene (OMIM# 142250)	52	55		
Entries with a fusion gene mutation	8	8		
Entries with a substitution mutation	899	1,031		
Entries with an insertion mutation	46	51		
Entries with a deletion mutation	116	150		
Hemoglobins with high oxygen affinity	79	90		
Unstable hemoglobins	121	134		
Methemoglobins	9	9		
Total hemoglobin variant entries	832	938		
Total thalassemia entries	336	383		
Total entries in both variant and thalassemia categories	43	48		
Total entries in database	1,125	1,273		

<sup>&</sup>lt;sup>a</sup>: Data from Hardison et al., 2002, <sup>b</sup>: Present report.

In addition to these changes in the operation of the query and history pages, there are several new features that have been added. Users can now query HbVar by gene location as well as by chain. For example, by selecting "Beta" and "Exon" the user will only get the mutations that are in the exons of the  $\beta$ -globin gene. Also under the "Mutation data" section there is a button to select regions by clicking on an image. There, a user can select a gene or a region between genes on the first image (Fig. 2A). When the user zooms in on a particular gene in order to select an exon or intron, the mutations documented in HbVar are displayed in a dense mode below the genes, grouped by category (substitutions, deletions, insertions, *etc.*) to give an impression of how much data is included in HbVar for each region (Fig. 2B).

In the "Occurrence" section, additional fields have been included to refine the query based on the population sample size tested. This is an important parameter, which is indicative of how representative the mutation frequencies are.

Automatic routines have been implemented to query for single nucleotide polymorphisms (SNPs) in the human  $\alpha$ - and  $\beta$ -globin loci via the University of California at Santa Cruz (UCSC) Genome Browser (Hinrichs et al., 2006). The user can choose to view the results either in a table format or as graphical display.

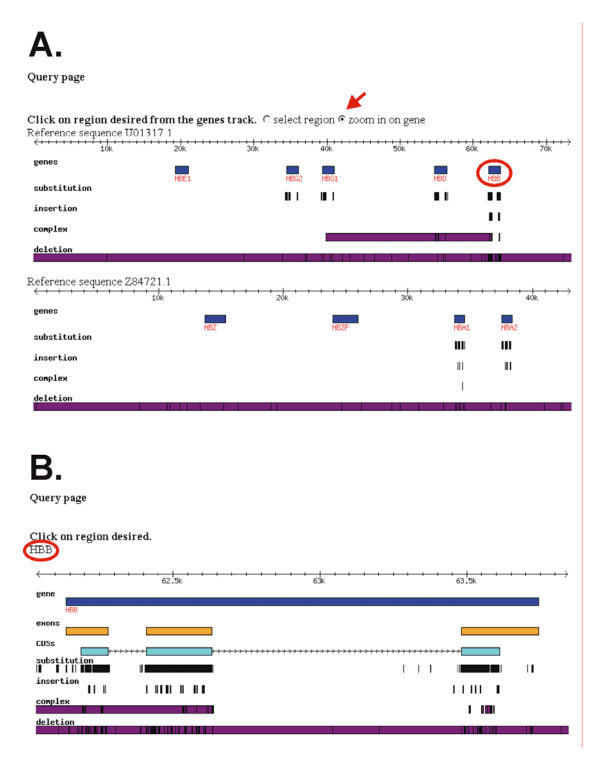


Figure 2. Overview of the new HbVar querying feature to select regions from an image. A: A user can select a particular region between genes or zoom in on a gene (indicated by the red arrow) to select a particular exon or intron. B: Outcome from selecting the human  $\beta$ -globin gene (HBB) region (circled in red). The various types of mutations included in HbVar (substitutions, insertions, deletions, and complex rearrangements) are displayed in a dense mode below the gene to give an indication of the amount of data currently available for each region. Mutant entries in these images are not "clickable."

#### DATABASE IMPLEMENTATION AND ACCESS

In its underlying implementation, HbVar was initially designed and built as a relational database, using Oracle as the database management system. We have recently switched from Oracle to PostgreSQL (http://www.postgresql.org), an object-oriented relational database management system (sometimes referred to as an object-relational database), which is available in several free and commercial versions. This choice was guided partly by the high costs of the Oracle license, compared to the free PostgreSQL system.

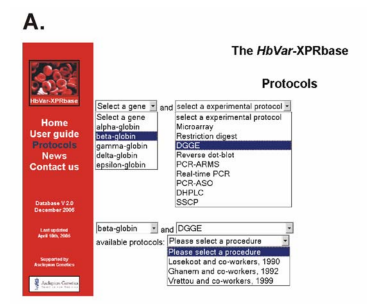
The HbVar database and associated resources at the Globin Gene Server (http://globin.bx.psu.edu, Hardison et al., 1994), such as the online *Syllabi*, are currently in use worldwide. Since January 2000, we recorded 15,859 accesses to the HbVar query page and 87,328 accesses to the online *Syllabi*, a significant increase compared to the previous years (Hardison et al., 2002, Patrinos et al., 2004). Users frequently contact the curators and the rest of the HbVar team members in order to submit new hemoglobin variants and/or thalassemia mutations, report missing information for existing mutants and pinpoint inconsistencies and/or erroneous entries. This is particularly important, since user input is vital in our effort to improve the data quality and accuracy. Therefore we encourage HbVar users to notify the curators regarding such issues (detailed contact information is available at the Globin Gene Server site).

#### XPRBASE, A HUMAN GLOBIN GENE MUTATION SCREENING PROTOCOLS DATABASE

We have constructed a separate database, named XPRbase (Fig. 3A), to provide a succinct summary listing of the protocols available for human globin gene mutation screening. The database currently contains 51 protocols for whole-gene, mutation-screening strategies (see also Patrinos et al., 2005a). These protocols are summarized in Table 2. XPRbase can be accessed on the World Wide Web at: http://www.goldenhelix.org/xprbase, and a link is provided from the HbVar homepage. Detailed instructions for both using and querying the database are also available from the same site. There has been no claim of ownership of the information stored in this database by anyone involved in this initiative. However, this compilation and representations of it are subject to copyright and usage principles to ensure that this resource remains freely available to all interested individuals.

**Table 2.** Summary of the Experimental Protocols Available in XPRbase to Screen for Sequence Alterations in the Human Globin Loci, Categorized by Gene and Mutation Detection Strategy

	Protocols	Globin genes				T 4 1	
Type		α	β	γ	δ	3	Total
	Restriction analysis	1	1	1	1	0	4
	Reverse dot-blot	1	3	0	0	0	4
'n ons	PCR-ASO	2	1	0	0	0	3
Known	PCR-ARMS	1	3	0	0	0	4
K	Real-time PCR	0	2	0	0	0	2
	GAP-PCR	8	0	1	1	0	10
	Microarrays	2	4	0	0	0	6
Unknown mutations	SSCP	3	2	0	0	1	6
	DGGE	1	3	2	1	0	7
	DG-DGGE	1	0	0	0	0	1
l II	DHPLC	1	3	0	0	0	4
Total		21	22	4	3	1	51



# В.

Primer	Sequence (5'>3')	Primer C (pmol)	Fragment name	Fragment size (bp)
A1	TTTAGTAGCAATTTGTACTGA	12.5	FR0	200
A2	GCCCTGGCTCCTGCCCTCCC	12.5		
GCA	CBCCCBCCCCGCCCGTGCCCCCGGGCCGCCCGCCCCGTACGGCTGTCATCACTTAGACCTCA	12.5	FR1	250
pCO4	CAACTTCATCCACGTTCACC	12.5		
apCO4	GGTGAACGTGGATGAAGTT	12.5	FR2	370
GC.B	CGCCCGCCCGCCCCGTGCCCCCGCGCCCCGCCCCCGCCCCCTGCAGCTTGTCACAGTGCAGCTCACT	12.5		
С	GTGTACACATATTGACCAAA	12.5	FR4	420
D	AGCACACAGACCAGCACGTT	12.5		
aD	AACGTGCTGGTCTGTGTGCT	12.5	FR5	300
X	ANATGCACTGACCTCCCACA	12.5		
pCO3	ACACAACTGTGTTCACTAGC	12.5	FR6	460
GC.B	As above	12.6		

Total volume: 50-uL reaction,

dNTPs: 200 uM of each dNTP,

MgCl2: 2.5 mM,

Taq DNA polymerase: 1 unit in supplied reaction buffer,

Genomic DNA: 0.5-1 ug,

0.05% W-1

Primers: Please see Table

Amplification conditions	Temperature	Time
1. Initial denaturation	94	4 min
2. Denaturation	94	1 min
3. Annealing	60	1 min 30 sec
4. Elongation	70	2 min
35 cycles to step 2		
5. Final elongation	72	7 min

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**Figure 3. A:** The protocols page from XPRbase. A user can select from two drop-down menus, which give the protocols that are available for mutation screening in every globin gene. When running the query "Find all DGGE protocols available for  $\beta$ -globin gene mutation screening," three protocols are displayed. **B:** Outcome of the query as described in A, consisting of the primer sequence, amplification conditions, reaction ingredients and the source reference(s), hyperlinked to PubMed, from where the publication(s) can be retrieved.

All database screens are based on the HTML language with some JavaScript and rely on Cascading Style Sheet (CSS) support. They are built using a custom-made PHP script that comprises the database's core engine, not only for menus and basic screens that display and parse files, but also for handling data querying. XPRbase is a flat-file database, derived from the *ETHNOS* V1.0 software, which facilitates the establishment of National Mutation Frequency databases (Patrinos et al., 2005b). A user guide provides some brief information on the operation and querying principles of XPRbase.

Database protocols can be added and/or modified only by the administrator through a dedicated administrator module (Sjozef van Baal and George Patrinos, unpublished). They are contained within an index file, namely "protocols.tab". When a new protocol is added to this file, a separate file named for the first author of the protocol is automatically created. This enables the administrator to enter the new protocol details by selecting this file from the protocol drop-down menu. To modify an existing protocol, the administrator only needs to select the desired protocol from the list and modify its contents in the designated area. All XPRbase data are freely available for academic research purposes, and data and accompanying descriptions are available for this website.

#### **FUTURE PROSPECTS**

During the 5 years that HbVar has been fully functional, it has become a key resource for information about sequence variation leading to hemoglobinopathies. One important feature is its constant update and improvement, mostly driven by the devotion of the researchers involved in this project and the valuable input from the end users, reporting erroneous entries and omissions and suggesting new additions. Given the increasing impact of HbVar on society and its growing content, we plan to expand the curator team by adding more expert advisors, and further to perform a user satisfaction survey to be able to identify weaknesses in the database contents and structure. The positive impact HbVar has on the globin research community is also illustrated by the fact that funding, whether dedicated or related to other projects, has always been available for keeping this resource alive, in an environment where funding opportunities for database development and curation are often limited and very hard to find (Patrinos and Brookes, 2005), frequently resulting in the discontinuation of many useful databases.

In order to ensure continuous data influx into HbVar, we plan to implement a broader search strategy that combines manual and electronic search procedures. This involves an expanded computerized search of all articles listed in the PubMed literature database (http://www.ncbi.nlm.nih.gov:80/entrez/query.fcgi?db=PubMed), in addition to the manual screening of a selection of core hematology journals. Also, in order to prevent globin gene mutation data from being overlooked, it is anticipated that HbVar will be more tightly linked to the journal *Hemoglobin*, following the successful recipe of the Waystation project (http://www.centralmutations.org) linked to the journal *Human Mutation*, and of the Human Gene Mutation Database (HGMD; http://www.hgmd.org, Stenson et al., 2003) currently linked to the journal *Human Genetics*. In addition, HbVar will be used as the main globin gene mutation data source for the portal currently under construction for the ITHANET project (http://www.ithanet.eu).

We also plan to enrich XPRbase, a vital component of the ITHANET portal, with additional experimental protocols and to create multiple links between individual HbVar entries and the protocols by which those mutations can be detected. Also, a graphical display of all oligonucleotide primers documented in XPRbase in relation to their respective genomic regions can improve the service provided by this database. Although this will require a more advanced technological platform, this feature would help in designing gap-PCR protocols to detect the over 40 known  $\delta\beta$ -thalassemia and Hereditary Persistence of Fetal Hb (HPFH) deletional mutants, provided that their deletion breakpoints are precisely mapped within HbVar. That is an ongoing effort, facilitated by the fact that the DNA sequence for the human  $\beta$ -globin gene cluster and surrounding regions is now complete. This will allow the precise identification of all deletion junctions and annotation of the DNA features affected by the deletions. Indeed, this information will be critical for interpreting all deletional mutants. Not only will the junction sequences allow better analysis and interpretation of the mutations, but they will also allow specialized screening strategies to be designed and implemented for each mutation.

The link between the HbVar and GALA databases (Giardine et al., 2003, Patrinos et al., 2004), coupled with the UCSC Genome Browser (Hinrichs et al., 2006), was the first step towards integrating the available resources at the Globin Gene Server (Hardison et al., 1994). A step further is PhenCode (Giardine et al., submitted), which connects the phenotype and clinical data in a variety of LSDBs, including HbVar and GenPhen, to the data on genome sequences, evolutionary history, and function available at the Genome Browser. Displaying locus-specific mutation data with clinical relevance on the Genome Browser makes it readily available to a wide audience, and facilitates the viewing of data from many sources in one context. On the other hand, links back to the original databases allow detailed queries within individual loci, which can then be further analyzed accordingly. This

combination of LSDBs and a powerful genome browser paves the way for drawing connections from genome to function to phenotype. Also, expansion of these connections to additional loci of clinical importance, perhaps building on the current WayStation project, will help in fulfilling the promise of the Human Genome Project to improve human health.

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