Recognition of fold- and function-specific sites in the ligand-binding domain of the thyroid hormone receptor-like family

Sonia Verma¹,²,³, Soumyananda Chakraborti¹, Om P. Singh¹, Veena Pande⁴, Rajnikant Dixit¹, Amit V. Pandey²,³* and Kailash C. Pandey¹,⁵*

¹Parasite-Host Biology Group, ICMR-National Institute of Malaria Research, New Delhi, India, ²Pediatric Endocrinology, Diabetology, and Metabolism, University Children’s Hospital, Bern, Switzerland, ³Translation Hormone Research Cluster, Department of Biomedical Research, University of Bern, Bern, Switzerland, ⁴Kumaun University, Nainital, Uttarakhand, India, ⁵Academy of Scientific and Innovative Research, Ghaziabad, Uttar Pradesh, India

Background: The thyroid hormone receptor-like (THR-like) family is the largest transcription factors family belonging to the nuclear receptor superfamily, which directly binds to DNA and regulates the gene expression and thereby controls various metabolic processes in a ligand-dependent manner. The THR-like family contains receptors THR, RARs, VDR, PPARs, RORs, Rev-erbs, CAR, PXR, LXR, and others. THR-like receptors are involved in many aspects of human health, including development, metabolism and homeostasis. Therefore, it is considered an important therapeutic target for various diseases such as osteoporosis, rickets, diabetes, etc.

Methods: In this study, we have performed an extensive sequence and structure analysis of the ligand-binding domain (LBD) of the THR-like family spanning multiple taxa. We have used different computational tools (information-theoretic measures; relative entropy) to predict the key residues responsible for fold and functional specificity in the LBD of the THR-like family. The MSA of THR-like LBDs was further used as input in conservation studies and phylogenetic clustering studies.

Results: Phylogenetic analysis of the LBD domain of THR-like proteins resulted in the clustering of eight subfamilies based on their sequence homology. The conservation analysis by relative entropy (RE) revealed that structurally important residues are conserved throughout the LBDs in the THR-like family. The multi-harmony conservation analysis further predicted specificity in determining residues in LBDs of THR-like subfamilies. Finally, fold and functional specificity determining residues (residues critical for ligand, DBD and coregulators binding) were mapped on the three-dimensional structure of thyroid hormone receptor protein. We then compiled a list of natural mutations in THR-like LBDs and mapped them along with fold and function-specific
mutations. Some of the mutations were found to have a link with severe diseases like hypothyroidism, rickets, obesity, lipodystrophy, epilepsy, etc.

**Conclusion:** Our study identifies fold and function-specific residues in THR-like LBDs. We believe that this study will be useful in exploring the role of these residues in the binding of different drugs, ligands, and protein-protein interaction among partner proteins. So this study might be helpful in the rational design of either ligands or receptors.

**KEYWORDS**
nuclear receptor, thyroid hormone receptor-like family, ligand-binding domain, sequence conservation, phylogeny, relative entropy, multi-harmony, functional specificity

**Background**

The nuclear receptor (NR) superfamily is the most abundant transcription factor (TFs) of metazoans (1, 2). NRs play an essential role in many biological processes such as development, metabolism, physiology, reproduction, and inflammation, as well as in many pathological processes such as cancer, diabetes, rheumatoid arthritis, neurologic and psychiatric syndromes, immunosuppression, hormone resistance syndromes, cardiovascular diseases, metabolic syndrome, etc. (3–8). Most NRs share a similar structural architecture organized into five or six modular regions (A to F), sharing variable degrees of homology (9, 10) [Figure 1A]. A brief description of the NR domains is as follows. A/B: N-terminal domain (NTD): The NTD is a highly disordered domain with little sequence conservation. The NTD contains the activator function-1 region (AF-1), which interacts with a variety of co-regulator proteins in a cell and promoter specific manner and is subject to alternative splicing and post-translational modifications (10–12). C: DNA binding domain (DBD): The C region is the highly conserved DNA binding domain (DBD) responsible for recognition and binding to specific DNA sequences called hormone response elements (HREs) in the promoter region of a target gene. The DBD is composed of two tetracysteine zinc-finger motifs (the N-terminal motif (ZF-1) and C-terminal motif (ZF-2)) exclusive to NRs and other sequence elements known as P-, D-, T- and A-boxes (Figure 1A) (13). These structural elements contribute to HRE specificity (P-box), dimerization (D-box), and contact with the DNA backbone (T- and A-boxes) (13). In the RXR-TR DBD structure, the T- and A-boxes are shown to directly interact with the minor groove of the spacer element (i.e 4 bp in TR) (Figure 1B) (14). The T and A-boxes also responsible for assisting DBDs to conduct domain-domain interactions on DNA (15). The C-terminal region of DBD contains the nuclear localization signal and is located in the (16) D: Hinge Region: The hinge region is a variable-length and poorly conserved linker region between the DBD and the LBD. Variability in the length of this region allows the C and E domains to adopt different conformations and consequently contributes to the selection of the DNA binding sites. E: Ligand-binding domain (LBD): LBD is less conserved than DBD. The LBD is an allosteric signaling domain that binds to ligands and also interacts directly with DBD and co-regulator proteins in the multi-domain structure of the receptor organized on its DNA response element (Figures 1D-F) (17, 18). This domain commonly contains 12 α-helices and four β-strands that fold into three parallel layers to form an alpha-helical sandwich and a hydrophobic ligand-binding pocket (LBP) at the base of the receptor (Figure 1C). The C terminus of LBD contains another activation function surface (AF-2), which is composed of helices 3, 4, and 12. Helix 12, is conformationally dynamic and stabilized upon ligand binding, and appears to communicate with AF-2 to facilitate interaction with different co-regulator proteins (19). F: The F region is not present in all NRs and its function remains unclear.

According to sequence alignment and phylogenetic analyses of the conserved C and E domains, NRs were classified into eight subfamilies (NR1 to NR8) (Table 1) (9, 20), which share the

Abbreviations: NR, nuclear receptor; THR-like, thyroid hormone receptor-like; TF, transcription factors; DNA, deoxyribonucleic acid; THR, thyroid hormone receptor; RXXR, retinoid X receptor; RAR, retinoic acid receptor; PPAR, peroxisome proliferator-activated receptor; VDR, vitamin D receptor; ROR, retinoic-related orphan receptor; CAR, constitutive androstane receptor; PXR, pregnane X receptor; LXR, liver X receptor; DBD, DNA-binding domain; LBD, ligand-binding domain; AF, activation function; HRE, hormone response element; LBP, ligand-binding pocket; DR, direct repeat; NCoR, nuclear receptor corepressor; MSA, multiple sequence alignment; HREs, hormone-response elements; DR, Direct repeat.
conserved modular structure (Figure 1A), except NR7 members who have two DBDs (9). Non-canonical NRs lack one of the two conserved domains (C in vertebrates and E in invertebrates) and are grouped into the NR0 subfamily (9, 20, 21).

Data on the origin of nuclear receptors suggest that they were not hormonal receptors with a high affinity for a particular ligand, and that feature was acquired later during evolution (22). The THR-like family evolved as receptors with selective ligands (3, 22). The thyroid hormone receptor-like (THR-like) family is one of the largest families belonging to the nuclear receptor superfamily. THR-like receptors are ligand-activated transcription factors that bind to the DNA and regulate gene
Like nuclear receptors, THR-like receptors are modular in structure and have the following functional domains: (1) N-terminal domain (A/B), (2) DBD, (3) flexible hinge region (D) (4) LBD, (E) (Figure 1A) (7). Among all the domains DBD is highly conserved, while the LBD is moderately conserved (3, 9, 23). The ligand-binding domain is the interaction site for ligands and co-regulators (co-activators (SRC1) or corepressors (NCoR, SMRT)) (24). Ligand binding in the binding pocket induces a conformational change in LBD, leading to the release of the corepressors and the recruitment of co-activators. This conformational change in LBD propagates to the DNA binding domain through direct contact that promotes the DBD to bind the DNA and the subsequent transactivation of genes (9, 25). Structural findings in multiple nearly full-length complexes have now shown that DBD and LBDs communicate, through direct contacts (13, 15, 17) (Figures 1D-F). THR-like receptors form monomers, homodimers and heterodimers, most commonly with RXR on bipartite HREs composed of tandem AGGTCA ‘half site’. The HREs for RAR, PPAR, and LXR differ from one another only by the number of base pairs between two hexameric direct repeats (DR1-D5) (Figures 1D-F) (9). The LBD is the main interaction site for ligands, coactivators, and corepressors and is also involved in the heteromerization of receptors. The ligands of THR-like receptors are small lipophilic molecules, including thyroid hormone, retinoic acid, fatty acids, heme, bile acids, and sterols. (Table 2). Based on sequence homology THR-like receptors are classified into 11 subfamilies i.e. (A) thyroid hormone receptor (THR) (B) retinoic acid receptor (RAR) (C) peroxisome proliferator-activated receptor (PPAR) (D) Rev-ErbA (E) E78C-like (F) RAR-related orphan receptor (RO) (G) CNR14-like (H) Liver X receptor-like (I) Vitamin D receptor-like (J) Hr96-like (K) Hr1-like (Table 2) (20).

THR-like receptors represent novel targets for the development of therapeutic agents for the treatment of numerous diseases, including type 2 diabetes, obesity dyslipidemia, atherosclerosis, and metabolic syndrome (7). There have been many recent advances in the development of new therapeutic agents for a subset of these receptors, including peroxisome proliferator-activated receptors, liver X receptors, and farnesoid X receptors (4, 6, 7). Considering the therapeutic potentials, we analyze the LBD of the THR-like family to predict the fold and function-specific sites by using information-theoretic methods such as relative entropy and sequence harmony. In the present study, we perform extensive in-silico analysis to retrieve essential details regarding phylogenetic relationships and functionally important residues in the LBD of THR-like receptors. Taken together, the results provide a necessary framework for structural and functional studies of these important families of protein. This study would be helpful in detail in understanding the molecular basis of selective mutation, which might open new possibilities for the design of a new class of therapeutic agents.

### Materials and methods

#### Data collection

To extract the protein sequences of the THR-like family, we searched various protein databases (NCBI, UniProt, EMBL) (https://www.ncbi.nlm.nih.gov/, https://www.uniprot.org/, https://www.embl.org/). We have created an HMM profile (Hidden Markov model) from the nuclear receptor family 1 (NR1) sequence seed alignment (26) (Figure 2). The HMM profile was searched against each protein database and full-length THR-like protein sequences were extracted using an internal script. CD-hit (Cluster Database at High Identity with Tolerance) was used to reduce sequence redundancy (27). Sequences that have more than 90% sequence similarities were removed from our database.

#### Mapping of ligand-binding domain and sequence extraction

THR-like family proteins have mainly two conserved domains DBD and LBD. To extract the LBD sequences from the full-length THR-like protein, we need to map the start and end of the domain in the sequence. To map the LBD in the protein sequences, we have designed a strategy. The THR-like LBD sequence seed alignment was downloaded from the conserved domain database (CDD) (28). Seed alignment was used to create an HMM profile using HMMER 3.0 (26). The HMM profile was searched for in THR-like full-length sequences. The Hmmsearch (26) output provides the start and end of the LBD in each sequence. The LBD sequences were extracted by using an in-house script (Figure 2).
Sequence and structure alignment

All THR-like LBD sequences were aligned using MAFFT with the default parameters (29) (Figure 2). In an evolutionary process, protein structures are more conserved than sequences. Therefore, to include the structure conservation information, we have created a structure sequence multiple sequence alignment of THR-like LBD. Structures of THR-like LBD protein belonging to different subfamilies (Table S1) were downloaded from the PDB database and aligned using the program STAMP (30), which is an integral tool in VMD (31). A structure-sequence MSA of THR-like LBD was created by aligning the structure profile and the sequence profile using the profile-profile alignment command in MAFFT (29). Jalview was to visualize and edit the MSA (32). The sequences of the CNR14-like subfamily (nematode), Hr96-like, and Hr1-like subfamily were

| S. No. | Subfamily group | Subfamily Name | Subfamily Members | Ligands | Functional role of the receptor |
|-------|----------------|---------------|-------------------|---------|--------------------------------|
| 1. A  | Thyroid hormone receptor | TRsa (Thyroid hormone receptor alpha) | Thyroid hormone | Regulation of metabolism, heart rate, and development of organisms |
| 2. B  | Retinoic acid receptor | RAR (Retinoic acid receptor α) | Vitamin A and related compounds | Cell differentiation, cell proliferation, and apoptosis |
| 3. C  | Peroxisome proliferator-activated receptor | PPARα (Peroxisome proliferator-activated receptor-α) | Fatty acids, prostaglandins | Cellular differentiation, development, and metabolism (carbohydrate, lipid, protein) and tumorigenesis |
| 4. D  | Rev-ErbA | Rev-ErbA | Heme | Regulators of clock gene |
| 5. E  | E78C-like (only found in arthropod, trematode, mollusk, nematode) | E78C (Ecdysone-induced protein 78C) | – | Development and regulate key processes during oogenesis. control ecdysone signaling in insects driving metamorphosis and molting |
| 6. F  | RAR-related orphan receptor | ROR (RAR-related orphan receptor-α) | Cholesterol, ATRA | Development of the cerebellum and lymph nodes, lipid metabolism, immune response, maintenance of bone. Circadian rhythm, bone metabolism, and retinal neurogenesis, Lymph node development and immune response, survival of T helper 17 cells |
| 7. G  | CNR14-like (nematode) | Sex-1 (Steroid hormone receptor csr/14) | – | – |
| 8. H  | Liver X receptor-like | E(78C (Ecdysone receptor, Ecr (arthropod)) | Ecdysteroids | regulators of cholesterol, fatty acid, and glucose homeostasis |
| 9. I  | Vitamin D receptor-like | VDR (vitamin D receptor) | Vitamin D | Mineral Metabolism |
| 10. J | Hr96-like | Hr96/Daf-12 (Nuclear hormone receptor HR96) | Cholestrol/dauchronic acid | Drug Metabolism and bilirubin clearance |
| 11. K | Hr1-like | Nuclear hormone receptor HR1 | – | – |
removed from the analysis due to low significance (short length, lower number). Motif analysis was carried out by using the MEME (Multiple EM for Motif Elicitation) suite (33). MEME searches for repeated, ungapped sequence patterns that occur in DNA or protein sequences for motif prediction.

**Phylogenetic analysis of THR-like ligand-binding domain**

A phylogenetic tree was generated from the alignment of multiple sequences of THR-like LBD using FastTree2 software with the default amino acid substitution model (34) (Figure 2). FastTree2 is known for its ability to handle large alignments and run speeds that are faster than other software. FastTree2 adds minimum-evolution subtree-pruning-regrafting (SPRs) and maximum-likelihood NNIs. FastTree2 uses heuristics to restrict the search for better trees and estimates a rate of evolution for each site. Further, the phylogenetic tree was visualized and analyzed by using Archaeopteryx software (35).

**Calculation of relative entropy (RE) scores of THR-like LBD**

Relative Entropy (RE) scores were calculated by comparing the amino acid probability distribution for each column of the multiple sequence alignment with that of the background distribution.

RE or the Kullback–Leibler divergence (KL divergence) was originally introduced by Solomon Kullback and Richard Leibler in 1951 as the direct divergence between two distributions (Kullback and Leibler 1951). It is used to measure the difference between an amino acid distribution $P$ and some background distribution $P_{null}$. The RE score of column $i$ is defined as

$$RE_i = \sum_{x=1}^{20} P_i(x) \log \frac{P_i(x)}{P_{null}(x)}$$

$P_i(x)$ is the probability of the occurrence of amino acid $x$ in column $i$ of the MSA, where $P_{null}$ is the background probability of amino acid $x$, which is generally calculated as the probability of finding an amino acid $x$ in all available protein sequences, i.e. protein sequences in Swiss-Prot database.

We used our in-house perl script to do the calculations (36, 37). The high-scoring residues were further mapped onto the thyroid hormone receptor alpha structure (PDB ID: 1NAV; Chain A) by carrying out profile-profile alignment between the sequence and structural alignment.

**Multi-harmony analysis**

Many protein families have subfamilies with functional specialization, such as the binding of different ligands or the participation of different protein-protein interactions. A small number of amino acids generally determine functional specificity. Multiple sequence alignments are often used to reveal functionally important residues within a protein family and identification of key residues that determine functional differences between protein subfamilies. We have used Multi-Harmony, an interactive web server for detecting sub-type-specific sites in LBD of THR-like receptors (38). The multi-Harmony server uses a combined method of sequence Harmony (SH) (39) and multi-Relief (MR) (40) to detect subfamily-specific residues. Sequence Harmony is based on Shannon’s entropy and determines to what extent amino acid compositions between groups differ. Multi-Relief identifies...
residues based on RELIEF, a state-of-the-art machine learning technique for feature weighting.

\[ SH_{A:B} = \sum_i p_i \log \frac{p_{A_i}}{p_{B_i} + p_{A_i}} \]

The SH score can be calculated as the relative entropy of group A relative to the sum of the probabilities of both groups \((p_A + p_B)\).

**Mutation analysis**

A collection of currently known natural-occurring THR-like LBD mutations was assembled from the UniProtKB database (41). The resulting dataset includes naturally occurring LBD mutations in the following receptors: PPAR, RAR, THR, LXR, VDR and ROR (Table S2). Based on the generated list, all mutations were classified according to our scoring scheme as fold and function-specific mutations. Furthermore, mutations were mapped to have an association with any disease in humans.

**Results**

**Retrieval and downstream filtering of sequences**

A total of 1276 THR-like receptor sequences were collected from the databases. We have used nuclear receptor family 1 (NR1) sequence seed alignment to search the database. Sequence redundancy was removed by CD-HIT software, keeping 90% sequence similarity as the clustering threshold. CD-hit provided 1176 sequence clusters. THR-like receptors consist of several domains out of them, two are major domains; DBD and LBD. As we were interested in studying LBD, we designed a strategy to fetch the LBD sequence out of a full-length THR-like sequence. We have created an HMM profile from the THR-like LBD sequence seed alignment downloaded from the conserved domain database. The HMM profile was searched against the THR-like full-length sequences database. We have used the bit score value \((X=114)\) as the threshold per domain in the hmmssearch command against the THR-like sequence database. The hmmssearch output also provides the start and end of the LBD in each sequence. The LBD sequences were extracted by using an in-house script. The short length (between 100aa and 130aa) LBD domain sequences were removed, and finally, 1158 THR-like LBD sequences were used for the study.

**Multiple sequence alignment of THR-like LBD**

Multiple sequence alignment of THR-like LBD sequences was performed using MAFFT software. To check the accuracy of alignment, we performed a profile-profile alignment. As structures are more conserved than sequences, we aligned the THR-like LBD structure profile created using STAMP software (10 known structures of THR-like LBD proteins; details in Table 2) with the THR-like LBD sequence profile. We employed WebLogo 3 (http://weblogo.threeplusone.com/create.cgi) (42) to visualize the conservation patterns that emerge from the profile-profile alignment. The WebLogo shown in Figure 3 represents the conserved residues in the LBD region of the THR proteins. In the WebLogo plot, all the hydrophobic residues (YVMEFIW) are colored black, while neutral amino acid (SGHTAP) and hydrophilic residues (RKDENQ) are colored green and blue, respectively (Figure 3). MEME Motif analysis provided 12 motifs in the protein sequences (Figure 3). These motifs of THR-like LBD provided information about seven highly conserved signaling motifs and five interaction sites. The interaction sites are the motifs where ligands, DBD and co-proteins interact. Motif A occupies consensus sequence positions 207-216. This is known as the inverse NR-box (LLxxL) motif and is one of the main ligand interaction sites. The NR box (NR interacting box LxxLL) plays an important role in the interaction of NRs with co-activators and therefore, in the regulation of transcription. Motif B occupies consensus sequence positions 222-232 and resides in a critical region for the co-activator function. Motif C also known as the LxxDDQ motif occupies positions 236-245 and is a region that is also critical to co-activator function. Motif D occupies positions 396-401 and plays a significant role in ligand binding. Motif E is an inverse NR-box (LLxxL) and occupies positions 455-459. Motif F occupies positions 498-504. Motif G occupies alignment positions 540-548. Motifs F and G are critical for coactivators binding.

**Phylogenetic analysis of THR-like LBD**

THR-like family is the largest ligand-activated nuclear receptor family, with members existing only in metazoans and not found in plants, algae, fungi, and protists. Fasttree2 (34) which shows good and fast performance with large alignments was used for the tree construction from the THR-like LBD multiple sequence alignment. LBD sequences are clustered based on sequence homology and lineage. The resultant monophyletic tree showed clustering of the sequences into 8 subfamilies (Figure 4). The clustering of the branches is from points near the center of the tree. Cluster 1 includes all thyroid hormone receptors (red in the phylogenetic tree). The second cluster includes all members of the retinoic acid subfamily (green in the phylogenetic tree). The third group is Vitamin D receptor-like subfamily (colored blue). The fourth cluster is The Liver X receptor-like subfamily. The fifth cluster is the peroxisome proliferator-activated receptor subfamily. The sixth cluster is The Rev-ErbA subfamily. The seventh cluster is The RAR-
related orphan receptor subfamily. The eighth cluster is The E78c subfamily from the phylogenetic tree analysis; it is predicted that there all these sequences are clustering based on their evolutionary relationships. Here is a brief description of the subfamilies.

Subfamily 1 - Thyroid hormone receptor: Thyroid hormone receptors are activated by binding to thyroid hormone. Thyroid hormone receptors play a critical role in the regulation of metabolism, heart rate, and development of organisms (3). TH receptors also have non-genomic effects leading to second messenger activation, and the corresponding cellular response (43). Thyroid hormone receptors regulate gene expression by binding to hormone response elements (HREs) in DNA as monomers, homodimers, or heterodimers with other nuclear receptors. The sequence logo shows conservation patterns that emerged from the multiple-sequence alignment of THR-like LBDs. The overall height of the stack represents the conservation of the sequence at a particular position, and the frequency of the respective amino acid at that position is indicated by the height of the amino acid symbol. The conserved motifs (red boxes) and interaction sites (site for ligand, coactivators/corepressors and DBD binding) are highlighted (green boxes).
receptors (retinoid X receptor (RXR)). There are two main classes of the thyroid hormone receptor, alpha (THRA) and beta (THRB). Mutations in THRs are associated with thyroid hormone resistance disease (44). Clinically mutations have been observed in both the THR isoforms, however, THRB gene mutations are much more common.

Subfamily 2- Retinoic acid: Retinoic acid receptors (RAR) are activated by all-trans retinoic acid and 9-cis retinoic acid. There are three retinoic acid receptors, RAR-alpha, RAR-beta, and RAR-gamma. RARs form heterodimers with members of the retinoid X receptor (RXR) subfamily (25). RARs also have nongenomic effects via activating kinase signaling pathways, which fine-tune the transcription of the RA target genes. The disruption of RA signaling pathways is thought to underlie the etiology of several hematological and non-hematological malignancies, including skin cancer, head/neck cancer, lung cancer, breast cancer, ovarian cancer, prostate cancer, renal cell carcinoma, pancreatic cancer, liver cancer, glioblastoma and neuroblastoma (45).

Subfamily 3- Vitamin D receptor-like: The VDR-like subfamily includes the vitamin D receptor, the pregnane X receptor, and the constitutive androstane receptor. Vitamin D receptor (VDR) heterodimerizes with retinoic acid upon binding calcitriol (an active form of vitamin D). The most potent natural agonist is calcitriol (1,25-dihydroxycholecalciferol) and vitamin D2 homolog ercalcitriol, 1-alpha,25-dihydroergocalciferol) is also a strong activator (3). Mutations in the VDR gene are associated with type II vitamin D-resistant rickets. A disorder of vitamin D metabolism resulting in severe rickets, alopecia, secondary hyperparathyroidism and hypocalcemia. The pregnane X receptor (PXR), is a nuclear receptor that senses the presence of foreign toxic substances and the metabolism (52).

Subfamily 4- Liver X receptor-like: The liver X receptors (LXR) are closely related to nuclear receptors such as the PPARs, FXR, and RXR. Liver X receptors (LXRs) are important regulators of fatty acid, cholesterol, and glucose homeostasis. Two LXR have been identified as LXRα and LXRβ. LXRα and LXRβ form heterodimers with the obligate partner retinoid X receptor (RXR), which is activated by 9-cis-13,14-dihydroretinoic acid (49). The LXR/RXR heterodimer can be activated by an LXR agonist (oxysterols) or an RXR agonist (9-cis-13,14-dihydroretinoic acid). Oxysterols, oxygenated derivatives of cholesterol, such as 22(R)-hydroxycholesterol, 24(S)-hydroxycholesterol, 27-hydroxycholesterol, and cholesterol acid, are natural ligands for LXR (49).

Subfamily 5- Peroxisome Proliferator-activated receptor: Peroxisome proliferator-activated receptors (PPARs) play essential roles in the regulation of cellular differentiation, development and metabolism (carbohydrate, lipid, protein) and tumor genesis of higher organisms. PPARs have three isoforms termed alpha, gamma, and delta (beta). PPARs heterodimerize with the retinoid X receptor (RXR) and regulate the transcription of targeted genes. Endogenous ligands for the PPARs include free fatty acids, eicosanoids, and Vitamin B3 (50).

Subfamily 6- Rev-ErbA: The Rev-Erb proteins are members of the nuclear receptor (NR) superfamily of intracellular transcription factors and key regulatory components of the circadian clock. There are two forms of the receptor, Rev-Erb alpha and Rev-Erb beta. These receptors act as key regulators of clock gene expression through transcriptional repression of Bmal1. Through their regulation of clock-controlled genes, the Rev-Erb proteins affect several physiological processes throughout the body, including metabolic, endocrine, and immune pathways (51).

Subfamily 7-RAR-related orphan receptor: The RAR-related orphan receptors (RORs) are intracellular transcription factors. There are three isoforms of ROR, ROR-α, -β, and -γ. The ROs are somewhat unique in that they appear to bind as monomers to hormone response elements as compared to other nuclear receptors which bind as dimers. ROs are activated by oxysterols. Some other natural substances have also been reported to bind to ROs, such as all-trans-retinoic acid binds with high affinity to ROR-β and -γ but not ROR-α. ROs act as lipid sensors and hence may play a role in the regulation of lipid metabolism (52).

Subfamily 8- Ecdysone-induced E78c: The members of the E78c protein induced by ecdysone are found in arthropods, trematodes, mollusks, and nematodes. E78c protein has been linked to insect development and metamorphosis (53). Ecdysone-induced protein 78C (E78), a nuclear hormone receptor closely related to Drosophila E75 and mammalian Rev-Erb and Peroxisome Proliferator-Activated Receptors was originally identified as an early ecdysone target. The previous study shows that E78C plays a role in oogenesis and is important for proper egg production and maternal control of early embryogenesis (53).

Prediction of fold-specific residues: Results of RE calculation

Fold-specific residues are responsible for maintaining the overall fold of the protein. Hence, such residues should be conserved across the THR protein alignment; an in-house Perl
script was used to predict the fold-specific residues by employing a relative entropy calculation (RE). RE calculations identify residues whose probability distribution is significantly different from their background probability distribution. RE, therefore, predicts residues that are not identified by the traditional scoring methods (36). RE scores calculated for each MSA alignment position were plotted against sequence conservation in WebLogo of THR-like LBD. High RE scores are represented by a sharp upward peak (blue color) in the graph at an alignment position (Figure 5). It was observed that the residues identified as fold-specific are scattered across the helices (Figures 6A, C). Interestingly, H4, H6, H8, H9, H10, and H11 contain certain residues with very high RE scores. To gain insight into the distribution of the high-scoring residues on a 3D structure, the top 30 residues with the highest scores were mapped onto the crystal structure of the thyroid hormone receptor alpha of homo sapiens (PDB ID: 1NAV). Table 3 summarizes the details of all residues which were indicated in the structure.

Prediction of function-specific residues: Results of multi-harmony calculation

Keeping in view the therapeutics of the THR-like family, we utilized another information-theoretic measure to explore distinct sequence signatures in the subfamilies. Though the proteins belonging to the THR-like family share the same protein fold, their ligand recognition and binding are different and very specific to each subfamily. It was of interest to explore the differentially conserved residues in one subfamily compared with another. The superposition of THR-like LBD structures reveals that the top part of the receptor is more or less similar compared to the bottom (which contains the LBP). This variability across THR-like receptors at the ligand-binding region allows THR-like receptors to recognize a diverse set of ligands. Therefore, to predict the specificity determining residues in the THR-like LBD we have used the multi-harmony web server. The Sequence Harmony (SH) and Multi-Relief (mR) methods in one web server allow simultaneous analysis and comparison of specificity residues. The SH scores calculated for each alignment position from the MSA were plotted against sequence conservation in weblogo of THR-like LBD (Figure 5, green color). The SH score lies between 0 and 1. The lower score shows the differential conservation of the residues at that alignment position. Refinements with low SH scores are the specificities that determine the residues. Low SH scores are represented by a downward peak (green color) in the graph at an alignment position (Figure 5). 30 residues with the minimum SH scores (considered 0.4 SH score as cutoff) are provided in Table 4. To get an idea of the distribution of the minimum SH scoring residues on a 3D structure, the top 30 residues with the minimum SH scores were mapped onto the homo sapiens crystal
structure of the thyroid hormone receptor alpha (PDB ID: 1NAV) (Figures 6B, D). The Result shows that most of the minimum SH scoring residues are around the ligand-binding cavity (Figure 6D). Table 4 summarizes the details of all residues which were indicated on the structure and a detailed list of residues with consensus sequences in all other subfamilies at that alignment position is provided in the supplementary information (Table S2).
In this study a list of all known natural mutations similar to THR-like LBD was composed from the UniProt knowledge database and the literature search (Table S3). Natural mutations were found in THRα, THRβ, RARα, RARβ, VDR, LXRα, PPARα, PPARγ and RORα, RORβ, and RORγ. Further mutations were classified according to our scoring scheme as fold and function specific (Table S3). THRA structure was used to map the predicted fold-, function-specific residues and natural mutations in THRA (Figures 7A–C). We have compiled a list of mutations that are associated with diseases in humans (Tables 5, 6). On the basis of mutation and the analysis of their results, it was observed that very highly conserved residues are not prone to mutations due to their essential role in protein function. The mutation in THRA LBD lies around the cavity (Figure 7C). We have observed that receptor-wise, THRβ has more mutation compared to THRA, VDR, and PPARα (Figure 8A). Mutations are present in all motifs and interaction sites but residues lying at motif G, motif A, motif C, motif F, motif B and interaction sites INS 5, INS 1, INS 2, INS 3 are more prone to mutations (Figure 8B). Mutational analysis also shows that most mutations are at the sites of functional significance compared to fold (Figure 8C).

### Discussion

In this study, we have combined both sequence and structural information from the THR-like LBDs into the multiple sequence alignment. Length-wise all THR-like

---

**TABLE 3** Residues with high Relative entropy (RE) scores. Residues are classified as fold-specific residues.

| S.No. | RE_score | Res name | Res Number |
|-------|----------|----------|------------|
| 1.    | 2.86093  | F        | 239        |
| 2.    | 2.736545 | F        | 232        |
| 3.    | 2.735418 | Q        | 247        |
| 4.    | 2.610201 | D        | 246        |
| 5.    | 2.39529  | K        | 234        |
| 6.    | 2.372856 | R        | 375        |
| 7.    | 2.334283 | L        | 302        |
| 8.    | 2.314478 | K        | 252        |
| 9.    | 2.293988 | Q        | 342        |
| 10.   | 2.222085 | H        | 381        |
| 11.   | 2.191814 | R        | 329        |
| 12.   | 2.183281 | E        | 257        |
| 13.   | 2.148886 | P        | 237        |
| 14.   | 2.095029 | D        | 328        |
| 15.   | 2.073641 | L        | 250        |
| 16.   | 2.023249 | E        | 315        |
| 17.   | 1.922814 | A        | 321        |
| 18.   | 1.875692 | L        | 318        |
| 19.   | 1.870668 | L        | 242        |
| 20.   | 1.77909  | F        | 300        |
| 21.   | 1.773385 | K        | 357        |
| 22.   | 1.748892 | L        | 251        |
| 23.   | 1.747452 | F        | 349        |
| 24.   | 1.702621 | P        | 399        |
| 25.   | 1.7008   | Y        | 267        |
| 26.   | 1.687514 | L        | 374        |
| 27.   | 1.672857 | E        | 403        |
| 28.   | 1.657511 | I        | 338        |
| 29.   | 1.648088 | F        | 215        |
| 30.   | 1.634578 | F        | 309        |

**TABLE 4** Residues with lower sequence harmony scores (SH). Residues are classified as function-specific residues.

| S.No. | SH_score | Res name | Res Number |
|-------|----------|----------|------------|
| 1.    | 0.20538  | C        | 392        |
| 2.    | 0.240834 | S        | 271        |
| 3.    | 0.258853 | K        | 220        |
| 4.    | 0.26986  | P        | 224        |
| 5.    | 0.29622  | L        | 292        |
| 6.    | 0.300949 | S        | 277        |
| 7.    | 0.301198 | A        | 281        |
| 8.    | 0.30187  | S        | 216        |
| 9.    | 0.307977 | M        | 280        |
| 10.   | 0.309991 | G        | 207        |
| 11.   | 0.313666 | A        | 263        |
| 12.   | 0.314308 | E        | 213        |
| 13.   | 0.316654 | R        | 228        |
| 14.   | 0.330136 | C        | 244        |
| 15.   | 0.332675 | S        | 240        |
| 16.   | 0.333702 | C        | 380        |
| 17.   | 0.334945 | M        | 259        |
| 18.   | 0.336499 | H        | 355        |
| 19.   | 0.341662 | I        | 222        |
| 20.   | 0.350112 | E        | 217        |
| 21.   | 0.351103 | V        | 265        |
| 22.   | 0.354899 | R        | 384        |
| 23.   | 0.359096 | C        | 255        |
| 24.   | 0.360052 | L        | 386        |
| 25.   | 0.36292  | S        | 296        |
| 26.   | 0.366648 | D        | 272        |
| 27.   | 0.366818 | Q        | 286        |
| 28.   | 0.369598 | L        | 396        |
| 29.   | 0.371785 | F        | 401        |
| 30.   | 0.372781 | D        | 211        |
ligand-binding domains are very similar. Finding conserved motifs should highlight regions of critical importance in ligand binding. Those regions can provide amino acid patterns that are a unique signature to the THR-like LBD. From the MSA we have found that the THR-like LBDs consist of seven conserved motifs and five interaction sites that are aligning with the previous studies (54). THR-like LBDs have the interaction site for the respective ligand and other co-proteins. It was observed that five interaction sites are present in the LBDs. Interaction site 1 lies within motif A, Interaction sites 2, 3, and 4 lie between motifs C and D, and interaction site 4 lies between motifs F & G.

The NR–boxes (LxxLL) or inverse NR–boxes (LLXXL) can bind to specific NR regions; they may have a role in the interaction between NRs, specifically in hetero- and homo-dimerization.

The MSA was used to draw a phylogenetic tree of the THR-like LBDs. The phylogenetic analysis performed depicts the distinct separation of THR-like LBDs into the monophyletic branches. The THR-like LBDs are clustered according to the lineage.

Fold- and function-specific residues found in any protein of interest, and these residues are differentially conserved among the subfamilies and provide them functional specificity, such as binding at different ligands or being involved in different protein-protein interactions. To predict fold-specific and function-specific residues in the THR-like family, we used information-theoretic methods. A high RE score at the alignment position corresponds to the fold-specific residue, and on the other hand, minimum SH score at the alignment position in MSA corresponds to the function-specific site. The structure of the THR-like LBD is composed of 12 α-helices folded as an antiparallel tri-layer alpha-helical sandwich fold in which antiparallel alpha-helices (H5, H8, H9; the “sandwich filling”) are flanked by two alpha-helices on one side (H2, H3) and three on the other (H6, H7, and H10; the “bread”) (9). The ligand-binding cavity is within the interior of the LBD and just below three antiparallel alpha-helical sandwich “fillings” (Figure 1C). The H12 is connected to H11 by a flexible loop which allows H12 to swing upon binding to a ligand, trapping the ligand and stabilizing the conformation of the NR (9). In this way, the LBD mediates recognition and interaction of ligands at the ligand-binding pocket (LBP), dimerization with LBDs from other NRs, interaction with co-regulators (coactivators [LxxLL motif in H4] or corepressors ([LxxxIxxxI/L motif in H1]) and ligand-dependent transactivation at the ligand-dependent activation function AF2 domain (H12), which enables the recruitment of co-regulators (21).

Additionally, to validate our study, we have compiled a list of natural mutations in THR-like LBDs and how they affect their function (Table S3). Since the interaction sites are mainly responsible for the ligand selectivity or co-proteins interactions, mutation analysis provides information that all the interaction sites and conserved motifs are prone to mutation (Figure 8B). Motif G and INS5 have high mutation count as compared to other motifs. Motif G and INS5 lie in helix H12 and H11, which play a crucial role in ligand trapping, binding, and stabilization of the receptors. Any mutation in these regions will lead to impairment of ligand binding, dimerization, and coreceptor binding.
### TABLE 5  THR-like receptors are involved in various diseases in humans.

| S.No. | Receptor | Involvement in disease |
|-------|----------|------------------------|
| 1.    | THRα     | Hypothyroidism, congenital, non-goitrous, 6 (CHNG6)  
   Disease description:  
   A disease characterized by growth retardation, developmental retardation, skeletal dysplasia, borderline low thyroxine levels, and high triiodothyronine levels. There is differential sensitivity to the action of thyroid hormones, with retention of hormone responsiveness on the hypothalamic-pituitary axis and liver but skeletal, gastrointestinal, and myocardial resistance. |
| 2.    | THRβ     | Thyroid hormone resistance, generalized, autosomal dominant (GRTHD)  
   Disease description:  
   An autosomal dominant disease characterized by high levels of circulating thyroid hormones (T3-T4), goiter, abnormal mental functions, increased susceptibility to infections, abnormal growth and bone maturation, tachycardia, and deafness. Affected individuals may also have attention deficit-hyperactivity disorders (ADHD) and language difficulties. Patients have normal or slightly elevated thyroid stimulating hormone (TSH). |
| 3.    | RARα     | Microphthalmia, syndromic, 12 (MCOPS12)  
   Disease description:  
   A form of microphthalmia, a disorder of eye formation that ranges from the small size of a single eye to the complete bilateral absence of ocular tissues (anophthalmia). In many cases, microphthalmia/anophthalmia occurs in association with syndromes that include non-ocular abnormalities. Patients with MCOPS12 exhibit variable features, including diaphragmatic hernia, pulmonary hypoplasia, and cardiac abnormalities. |
| 4.    | RARβ     | Intellectual developmental disorder with or without epilepsy or cerebellar ataxia (IDDECA)  
   Disease description:  
   An autosomal dominant neurodevelopmental disorder that manifests with variable features of mild to severe intellectual disability, developmental delay, autism spectrum disorder, cerebellar ataxia, and epilepsy. |
| 5.    | VDR      | Selective pituitary thyroid hormone resistance (PRTH)  
   Disease description:  
   Variant form of thyroid hormone resistance and is characterized by clinical hyperthyroidism, with elevated free thyroid hormones but inappropriate normal serum TSH. Unlike GRTH, where the syndrome usually segregates with a dominant allele, the mode of inheritance in PRTH has not been established. |
| 6.    | PPARY    | Obesity  
   Disease description:  
   A condition characterized by an increased body weight beyond the limitation of skeletal and physical requirements as the result of excessive accumulation of body fat. |
| 7.    | RORα     | Rickets vitamin D-dependent 2A (VDDR2A)  
   Disease description:  
   A disorder of vitamin D metabolism results in severe rickets, hypocalcemia, and secondary hyperparathyroidism. Most patients have total alopecia in addition to rickets. |
| 8.    | RORβ     | Lipodystrophy, familial partial, 3 (FPLD3)  
   Disease description:  
   A form of lipodystrophy characterized by marked loss of subcutaneous fat from the extremities. Facial adipose tissue may be increased, decreased, or normal. Affected individuals show an increased preponderance of insulin resistance, diabetes mellitus, and dyslipidemia. PPARG defects can lead to type 2 insulin-resistant diabetes and hypertension. PPARG mutations may be associated with colon cancer. |
| 9.    | Glioma 1 (GLM1) | Glioma 1 (GLM1)  
   Disease description:  
   Gliomas are benign or malignant central nervous system neoplasms derived from glial cells. They comprise multiforme astrocytomas and glioblastomas derived from astrocytes, oligodendrogliomas derived from oligodendrocytes, and ependymomas derived from ependymocytes. |
| 10.   | Polymorphism | Genetic variations in PPARG define the quantitative trait locus 1 of the body mass index (BMIQ1). The body mass index (BMI) reflects the amount of fat, lean mass, and body build. Genetic variations in PPARG influence the carotid intimal medial thickness (CIMT). CIMT is a measure of atherosclerosis that is independently associated with traditional atherothrombotic cardiovascular disease risk factors and coronary atherosclerotic burden. 35 to 45% of the variability in multivariate-adjusted CIMT is explained by genetic factors. |

(Continued)
TABLE 5 Continued

| S.No. | Receptor | Involvement in disease |
|-------|----------|------------------------|
| 9.    | RORγ     | **Disruption phenotype**<br>Mice show decreased adipocyte size and high insulin sensitivity, leading to improved control of circulating fatty acids. Mutants are protected against hyperglycemia and insulin resistance in the state of obesity. Loss of circadian pattern of some clock genes expression in peripheral tissues and massive apoptosis of thymocytes. Knockout mice for isoform 2 lack all lymph nodes and Peyer patches, as well as LTI cells. They also show a reduction of T(H)17 cells in the lamina propria by at least 10 times to less than 1% of T (H) cells. Mice are less susceptible to autoimmune inflammatory diseases. |

TABLE 6 List of THR-like receptor LBD mutations associated with diseases in humans.

| S. No. | Receptor | Mutation | Classification | Alignment Position | Disease-Associated |
|--------|----------|----------|----------------|--------------------|-------------------|
| 1      | THRα     | N359Y    | Function       | 484 in CHNG6; atypical phenotype; weak reduction in transcriptional activation |
| 2      | THRα     | A263S    | Function       | 258 in CHNG6, no effect on T3 binding; no effect on thyroid hormone-dependent transcriptional activation |
| 3      | THRβ     | R429Q/W  | Fold           | 506 in PRTH |
| 4      | THRβ     | S360L    | Function       | 368 in GRTHD |
| 5      | THRβ     | V349M    | Fold           | 367 in GRTHD |
| 6      | THRβ     | V348E    | Function       | 366 in GRTHD |
| 7      | THRβ     | G347E/A  | Fold           | 364 in GRTHD |
| 8      | THRβ     | L346F    | Fold/Function  | 363 in GRTHD |
| 9      | THRβ     | G345R/V/S| Fold           | 361 in GRTHD |
| 10     | THRβ     | G344E/A  | Function       | 360 in GRTHD |
| 11     | THRβ     | K342I    | Function       | 343 in GRTHD |
| 12     | THRβ     | L341P    | Function       | 336 in GRTHD |
| 13     | THRβ     | Q340H    | Function       | 335 in GRTHD |
| 14     | THRβ     | R338W/L  | Fold           | 295 in GRTHD |
| 15     | THRβ     | Del337T  | Function       | 294 in GRTHD |
| 16     | THRβ     | A335P    | Function       | 292 in GRTHD |
| 17     | THRβ     | E333D    | Fold           | 277 in GRTHD |
| 18     | THRβ     | G332R/E  | Function       | 276 in GRTHD |
| 19     | THRβ     | N331D    | Function       | 275 in GRTHD |
| 20     | THRβ     | L330S    | Fold           | 273 in GRTHD |
| 21     | THRβ     | T329I    | Function       | 272 in GRTHD |
| 22     | THRβ     | T327A    | Function       | 269 in GRTHD |
| 23     | THRβ     | Y321C/H  | Fold           | 262 in GRTHD |
| 24     | THRβ     | R320H/G  | Function       | 261 in GRTHD |
| 25     | THRβ     | A317T/S  | Function       | 258 in GRTHD; impairs hormone binding |
| 26     | THRβ     | R316H/C  | Fold           | 257 in PRTH; impairs hormone binding |
| 27     | THRβ     | A268G    | Function       | 207 in GRTHD |
| 28     | THRβ     | R243W    | Fold           | 80 in GRTHD |
| 29     | THRβ     | A234T    | Function       | 71 in GRTHD; impairs hormone binding and ligand-dependent conformational changes |
| 30     | RARβ     | R394C    | Function       | 523 in MCO12; increased transcriptional response to retinoic acid ligands |
| 31     | RARβ     | R394S    | Function       | 523 in MCO12; increases transcriptional response to retinoic acid |
| 32     | RARβ     | G303A    | Fold           | 361 in MCO12; Increased transcriptional response to retinoic acid ligands |
| 33     | RARβ     | L220P    | Function       | 204 in MCO12; Increased transcriptional response to retinoic acid ligands |

(Continued)
were further classified into fold- and function-specific residues according to our scoring scheme and analyzed how the mutation affects the function of the protein. Most of the mutations are function-specific which signifies that mutations are linked to impairment in ligand and co-receptor binding that leads to malfunctioning of the receptor and is associated with diseases in humans (55–60). We have found that 47 THR-like LBD mutations are critical and linked to diseases in humans (Table 7). Mutations in the LBD of THR\(\alpha\), THR\(\beta\), RAR\(\alpha\), RAR\(\beta\), VDR, PPAR\(\gamma\), ROR\(\alpha\), ROR\(\beta\), and ROR\(\gamma\) receptors are associated with several diseases (Table 6). A brief description of the diseases is provided for reference (Table 5). Among all receptors, THR\(\beta\) LBD is more prone to mutation and is involved in GRTHD, GRTHR, and PRTH diseases. THR\(\beta\) mutations A317T/S and A234T are involved in GRTHD and affect hormone binding, THR\(\beta\) mutation R316H/C is involved in PRTH and also impairs hormone binding (Table 6). THR\(\beta\) residues R316 and A317 constitute the ligand binding domain. The A317 side chain interacts with iodine, sculpts the T3 in the ligand-binding pocket, and R316 is a part of the polar cluster in THR\(\beta\). Both mutants show decreased T3 affinity and weak transcription activity. The mutant R316H fails to dimerize which explains the weak transcription (61). The residue R316 forms two hydrogen bonds with the helix1, upon mutation to H316 both the hydrogen bond is lost which leads to displacement of helix1 and thereby loop after helix1. The stability of helix1 is required for ligand binding and dimerization (61). The mutant A317T repositions 3,5,3’-triiodothyroacetic acid, distending the face of the receptor that binds the coregulators and (61). Mutation analysis has also shown that some alignment positions are very critical (Table 7). Two and more than two receptors having mutations at these positions lead to impairment of LBD function resulting in the occurrence of disease. Three receptors have the mutation, THR\(\beta\) (L346F), VDR (H305Q) and PPAR\(\gamma\) (F388L) at alignment position 363 and are associated with GRTHD, VDDR2A and FPLD3 diseases, respectively (55, 57, 62). These findings suggest that the spatial position of the residues is also important with the residue (polar, nonpolar, and neutral) at that position and may help in the selectivity, binding and interaction of ligands with the coactivators and co-repressors.

Conclusions

In conclusion, THR-like receptors are vital transcription regulators and therefore mentioned information acquired can be used in a variety of ways. Phylogenetic analysis also provided
new insights for THR-like clustering and identified several key regions that exist through evolution in the NR LBD. The mutation analysis highlighted mutational hotspots while also providing insights into their effects, especially when they are localized in conserved motifs of THR-like LBD. Due to the diverse array of genes regulated by these proteins, along with the fact that many drugs are not explicitly specified for one receptor, drugs that target THR-like receptors tend to have unwanted side effects. Improving our understanding of THR-like receptors could pave the way for future therapeutics. Our study might be helpful to pharmaceutical companies and academic researchers in developing synthetic ligands that can specifically target these receptors. The conserved motifs and interaction sites can be used as selected targeted regions for rational design and discovery of novel drugs. The development of new drugs can be achieved through specific in silico techniques that can compose...
ligands, which can interact with specific conserved regions and force new alterations in the protein’s dynamics.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding authors.

Author contributions

All authors contributed to the implementation of the research. SV, KP, and AP conceived and designed the study. SV, SC, KP, AP, OS, VP, and RD wrote the paper. All authors contributed to the article and approved the submitted version.

Funding

SV was supported by the Swiss Government Excellence Scholarship award number 2017.0601 (2017–2018) and ICMR-SRF (ISRM/11(98)/2019) fellowship during the study. Open access funding was provided by the University of Bern and swissuniversities. Open access funding was provided by the University of Bern.

Acknowledgments

The authors thank ICMR-NIMR and University Children’s Hospital Bern, University of Bern, Bern, Switzerland for their computational and infrastructure facility. SV would like to acknowledge Professor Andrew M Lynn, SC&IS, JNU for providing the scripts used in the study. SV also would like to thank Dr. Poonam Vishwakarma for their critical suggestion during the work. SC thankfully acknowledges the Ramalingaswami Fellowship (BT/RLF/Re-entry/09/2019).

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher’s note

All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fendo.2022.981090/full#supplementary-material
in response to phenobarbital. Mol Pharmacol (2002) 61(1):1–6. doi: 10.1124/mol.61.1.1

49. Willy PJ, Umeseo K, Ong ES, Evans RM, Heyman RA, Mangelsdorf DJ, et al. LXR, a nuclear receptor that defines a distinct retinoid response pathway. Genes Dev (1995) 9(9):1033–45. doi: 10.1101/gad.9.9.1033

50. Michalk L, Auehrs J, Berger JP, Chatterjee VK, Glass CK, Gonzales PJ, et al. International union of pharmacology. LXI. Peroxisome proliferator-activated receptors. Pharmacol Rev (2006) 58(4):726–41. doi: 10.1124/pr.58.4.5

51. Dumas B, Harding HP, Choi HS, Lehmann KA, Chung M, Lazar MA, et al. A new orphan member of the nuclear hormone receptor superfamily closely related to Rev-erb. Mol Endocrinol (1994) 8(8):996–1005. doi: 10.1210/mend.8.8.7997240.

52. Solt LA, Burris TP. Action of RORs and their ligands in (patho)physiology. Trends Endocrinol Metab (2012) 23(12):619–27. doi: 10.1016/j.tem.2012.05.012

53. Uryu O, Ameku T, Niwa R. Recent progress in understanding the role of ecdysteroids in adult insects: Germline development and circadian clock in the fruit fly, Drosophila melanogaster. Zool Lett (2015) 1:32. doi: 10.1186/s40851-015-0035-2

54. Mina T, Papageorgiou L, Ethimiadou A, Bacopoulou F, Viachakis D, Chrousos GP, et al. A comprehensive structural and functional analysis of the ligand-binding domain of the nuclear receptor superfamily reveals highly conserved signaling motifs and two distinct canonical forms through evolution. World Acad Sci J (2020) 1:264–74. doi: 10.3892/wasj.2020.30

55. Huber BR, Desclaux MA, West BL, Cunha-Lima ST, Nguyen HT, Baxter JD, et al. Thyroid hormone receptor-beta mutations conferring hormone resistance and reduced corepressor release exhibit decreased stability in the n-terminal ligand-binding domain. Mol Endocrinol (2003) 17(1):107–16. doi: 10.1210/me.2002-0097

56. Behr M, Loos U. A point mutation (Ala229 to thr) in the hinge domain of the c-erbA beta thyroid hormone receptor gene in a family with generalized thyroid hormone resistance. Mol Endocrinol (1992) 6(7):1119–26. doi: 10.1210/mend.6.7.1324420.

57. Hegele RA, Cao H, Frankowski ST, Leff T. PPARG F388L, a transactivation-deficient mutant, in familial partial lipodystrophy. Diabetes (2002) 51(12):3586–90. doi: 10.2337/diabetes.51.12.3586

58. Surol M, Chitayat D, Caron V, Chassaing N, Bitoun P, Patry L, et al. Recessive and dominant mutations in retinoic acid receptor beta in cases with microphthalmia and diaphragmatic hernia. Am J Hum Genet (2013) 93(4):765–72. doi: 10.1016/j.ajhg.2013.08.014

59. Guissart C, Latypova X, Rollier P, Khan TN, Stamberger H, McWalter K, et al. Dual molecular effects of dominant RORA mutations cause two variants of syndromic intellectual disability with either autism or cerebellar ataxia. Am J Hum Genet (2018) 102(5):744–59. doi: 10.1016/j.ajhg.2018.02.021

60. Kristjansson K, Rut AR, Hewison M, O’Riordan JL, Hughes MR. Two mutations in the hormone binding domain of the vitamin D receptor cause tissue resistance to 1,25-dihydroxyvitamin D3. J Clin Invest (1993) 92(1):12–6. doi: 10.1172/JCI16539

61. Huber BR, Sandler B, West BL, Cunha Lima ST, Nguyen HT, Apriletti JW, et al. Two resistance to thyroid hormone mutants with impaired hormone binding. Mol Endocrinol (2003) 17(4):643–52. doi: 10.1210/me.2002-0095

62. Arita K, Nanda A, Wessagowit V, Akiyama M, Alsaleh QA, McGrath JA, et al. A novel mutation in the VDR gene in hereditary vitamin D-resistant rickets. Br J Dermatol (2008) 158(1):168–71. doi: 10.1111/j.1365-2133.2007.08232.x.