Uncovering new disease indications for G-protein coupled receptors and their endogenous ligands

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Abstract

Background: The Open Targets Platform integrates different data sources in order to facilitate identification of potential therapeutic drug targets to treat human diseases. It currently provides evidence for nearly 2.6 million potential target-disease pairs. G-protein coupled receptors are a drug target class of high interest because of the number of successful drugs being developed against them over many years. Here we describe a systematic approach utilizing the Open Targets Platform data to uncover and prioritize potential new disease indications for the G-protein coupled receptors and their ligands.

Results: Utilizing the data available in the Open Targets platform, potential G-protein coupled receptor and endogenous ligand disease association pairs were systematically identified. Intriguing examples such as GPR35 for inflammatory bowel disease and CXCR4 for viral infection are used as illustrations of how a systematic approach can aid in the prioritization of interesting drug discovery hypotheses. Combining evidences for G-protein coupled receptors and their corresponding endogenous peptidergic ligands increases confidence and provides supportive evidence for potential new target-disease hypotheses. Comparing such hypotheses to the global pharma drug discovery pipeline to validate the approach showed that more than 93% of G-protein coupled receptor-disease pairs with a high overall Open Targets score involved receptors with an existing drug discovery program.

Conclusions: The Open Targets gene-disease score can be used to prioritize potential G-protein coupled receptors-indication hypotheses. In addition, availability of multiple different evidence types markedly increases confidence as does combining evidence from known receptor-ligand pairs. Comparing the top-ranked hypotheses to the current global pharma pipeline serves validation of our approach and identifies and prioritizes new therapeutic opportunities.

Keywords: G-protein coupled receptors, Drug discovery, Data integration, Target identification

Background

There are currently 827 known human G-protein coupled receptors (GPCRs) of which 406 are non-olfactory [1]. Together, this amounts to approximately 2% of all known protein-coding genes. They are, however, the largest ‘target’ class of the ‘druggable genome’ representing approximately 19% of the currently available drug targets [2, 3]. They have long played a prominent role in drug discovery [4] – so much so, that as of this writing, 475 FDA approved drugs act on GPCRs [5]. Several reasons account for this over-representation. GPCRs have ligand binding sites on the outer cell surface membrane, and potent effects can be achieved even from small ligand concentrations [2]. Some, but not all GPCRs have endogenous peptidergic ligands, small proteins produced by other cells that bind to the GPCR and trigger the downstream signalling cascade. Thus, endogenous peptides also provide a good starting point for the design of potential new drug targets due to their high tractability, specificity, safety, tolerability, and efficacy, as well as lower production complexity than other biopharmaceuticals [6]. These characteristics make GPCRs and their endogenous peptidergic ligands an extremely promising category of drug targets to investigate [7, 8].

To link potential drug targets, such as GPCRs, to disease indications, several public databases integrating various types of evidence are available including PHAROS [9], DisGeNET [10], The Monarch Initiative [11], and DISEASES [12] as well as the recently developed Open Targets...
platform [13]. This public-private platform integrates a large number of different data sources to provide evidence supporting the association between genes which could be known or new potential drug targets and human diseases [13]. As of October 2017, the Open Targets platform covers more than 26,000 genes which include both protein-coding as well non-coding gene identifiers and 9150 disease and phenotypic terms. In total, it consolidates evidence for nearly 2.6 million potential target-disease pairs. A scoring scheme was developed capturing the overall confidence and strength of a target-disease association given the available evidence such that the resulting association score ranging from 0 (“no evidence”) to 1 (“strongest evidence”) combines the observation frequency, the magnitude or strength, and the confidence in the source of evidence for a given target-disease association [13].

This is an exceedingly large number of hypotheses to analyse which raises the question of how a drug discovery scientist might prioritize amongst them. Potential ranking strategies might include the overall Open Targets score, the number of different types of evidence supporting the hypothesis, or other measures computed over the Open Targets database, such as mutual information [14] or machine learning approaches [15], that relate a given target-disease pair to other, similar hypotheses. Criteria to consider that are highly relevant to drug discovery but may currently reside outside the scope of the Open Targets platform include, for example, disease incidence and prevalence, unmet medical need, the availability of disease models and biomarkers [16], and druggability of the target [17].

Here we hypothesize that the evidence collected in the Open Targets platform supporting gene-disease associations can be used effectively to identify and prioritize target hypotheses for drug discovery. Focussing on a protein class of particular interest to drug discovery as a use case, we outline an innovative approach to identify and prioritize potential new GPCR and endogenous peptidergic therapeutic targets using the data behind the target-disease pairs from the Open Targets platform. First, we describe the distribution of the target-disease pairs and corresponding scores in the Open Targets platform database. Then we identify and characterize sets of GPCRs as well as their endogenous peptidergic ligands in the context of the Open Targets platform. Lastly, we compare the top-ranked GPCR and peptidergic targets to the current global pharma pipeline to validate our approach and to identify potential new disease indications and therapeutic opportunities.

Results
Distribution of the overall Open Targets score and relationship to individual data types
At the time of this analysis the Open Targets Platform integrates fifteen different data sources organized into seven different data types: genetic association, somatic mutations, RNA expression, known drug targets, affected molecular pathways, animal models, and text mining [13]. Each gene-disease pair receives a set of scores, each ranging from 0 to 1, representing the seven different data types as well as an overall cumulative score. These scores are designed to incorporate measures of the frequency, effect size, and confidence of the observed gene-disease evidence [13]. To examine the distribution of the resulting scores we plotted the empirical density and the cumulative distribution of the overall score, respectively (Fig. 1a). These plots suggest a mixture of distributions with most gene-disease pairs receiving scores near zero (median overall score = 0.057), a relatively broad peak around 0.15 and two other peaks around 0.55 and 1, respectively. Interestingly, the 95th percentile of the overall score is approximately 0.5 and approximately 2.7% of the gene-disease pairs in Open Targets have the maximum score of 1.

As has been observed elsewhere [18], the number of gene-disease pairs with a positive score varies considerably between the different data types. For example, 46% of the pairs had a literature mining score greater than zero compared to less than 5% of pairs with somatic mutation, known drugs, or affected pathways scores greater than zero, respectively (Fig. 1b). While the Open Targets Platform integrates many different data sources, most disease-gene associations (97%) are supported by only one or two different data sources and only a fraction of pairs (0.44%) have evidence from 4 or more different types of data sources. Comparing the overall score of a disease-gene association against the number of data types where the individual data type score is positive shows that the more independent data sources supporting a given gene-disease association the higher the overall score (Fig. 1c).

Characterizing GPCRs and endogenous ligands
We obtained a list of 403 human G-protein coupled receptors (GPCRs) from IUPHAR [19] of which 397 mapped to unique Entrez gene identifiers. In addition, from the same source, we obtained a list of 529 human endogenous ligands [19] which mapped to 412 unique Entrez gene identifiers. It should be noted that some genes encode multiple different peptides (e.g. the GCG gene encodes glucagon, GLP-1, and GLP-2). 119 of the GPCRs and 127 of the endogenous ligands are known to interact according to IUPHAR [19] forming 681 unique receptor-ligand pairs at the gene level. Of these pairs, 34 are 1:1 relationships meaning a GPCR binds exactly one endogenous ligand and vice versa. The remaining GPCR-endogenous ligand pairs are comprised of GPCRs which have up to 17 ligands (Fig. 2a) and ligands that interact with up to 8 different GPCRs (Fig. 2b).

Both GPCRs and endogenous ligands have a considerably higher number of associated disease terms in Open Targets
than other classes of genes. The average number of diseases associated with GPCRs is 198 and 413 for endogenous ligands while the average number of associated diseases for all other genes is 119 which is statistically significantly lower \( p = 1.8 \times 10^{-34} \) and \( p = 5.5 \times 10^{-117} \), respectively, Wilcoxon rank sum test. Interestingly, the number of associated disease terms is actually lower than expected for endogenous ligands when we use the relatively stringent 0.5 threshold for the overall Open Targets score but remains higher than expected for GPCRs (23 and 32, respectively, compared to the average of 26) but these comparisons are not significant at the 5% level when using the Wilcoxon rank sum test \( p = 8.3 \times 10^{-2} \) and \( p = 8.0 \times 10^{-1} \), respectively) (Fig. 2c). The increased number of disease associations for GPCRs and endogenous ligands is also reflected in the overall distribution of the Open Targets association score (Fig. 2d).

**Combining GPCR and endogenous ligand disease association evidences**

Known GPCR-endogenous ligand pairs can be used to accumulate additional evidence supporting a particular disease hypothesis of interest. For example, the evidence collected in the Open Targets platform suggests that galanin, an endogenous ligand for the GPCR galanin receptor type 2 (GALR2), plays an important role in epilepsy, one of the most common neurological disorders (overall score = 1.0). Indeed, galanin has long been suggested as a potential target to treat epilepsy [20]. In particular, there is evidence found through literature mining (score = 0.004) indicating that galanin depletion from the hippocampus may contribute to the maintenance of seizure activity [21], as well as genetic evidence (score = 1.0) showing that a galanin loss-of-function mutation leads to epilepsy in humans [22]. Interestingly, a recent paper suggests GALR2 as a more suitable potential drug target to treat epilepsy [23], but this literature mining result is currently the only type of evidence supporting the GALR2-epilepsy association in the Open Targets Platform. As a result, the corresponding overall score is a relatively low 0.018 which corresponds to the 20th percentile (Fig. 1a) and by itself does not stand out as a compelling new therapeutic target hypothesis. However, viewing the latter evidence together with the strong...
genetic evidence for galanin leads to a much stronger hypothesis. As this example illustrates, it may be advantageous to consider ligand-receptor pairs in concert to develop new hypotheses. Additional examples highlighting GPCR-ligand pairs of interest are listed in Table 1.

To more systematically identify potential disease indications associated with both an endogenous GPCR ligand and its receptor, we assembled GPCRs and their corresponding endogenous ligands that shared the same disease associations in the Open Targets Platform.

Figure 3a and the corresponding Additional file 1: Table S1 shows the overall Open Targets score for

### Table 1: Examples of known GPCR-endogenous ligand pairs with matching disease indications and corresponding Open Targets overall scores

| Disease                | GPCR Name | Score | Endogenous ligand Name | Score |
|------------------------|-----------|-------|-------------------------|-------|
| Epilepsy               | GALR2     | 0.02  | GAL                     | 1.00  |
| Obesity                | MC1R      | 0.04  | POMC                    | 1.00  |
| Alzheimer's disease    | FPR2      | 0.04  | APP                     | 1.00  |
| Inflammatory bowel disease | CCR3   | 0.05  | CCL7                    | 0.87  |
| Hypertension           | AGTR2     | 0.15  | AGT                     | 1.00  |
| Rheumatoid arthritis   | CCR6      | 0.99  | CCL20                   | 0.06  |
| Macular degeneration   | CX3CR1    | 1.00  | CX3CL1                  | 0.05  |
| Biliary dyskinesia     | SCTR      | 1.00  | VIP                     | 0.02  |
| osteoporosis           | CALCR     | 1.00  | ADM                     | 0.04  |
| vascular disease       | EDNRA     | 1.00  | EDN2                    | 0.04  |
disease-GPCR pairs plotted against the score for corresponding disease-endogenous ligand pairs. For this analysis, pairs without any evidence were assigned a score of 0. If there was a strong correlation between disease-GPCR pairs and pairs of the same disease and corresponding endogenous ligand, we would expect most disease-gene pairs in this plot scatter around the diagonal. However, the observed correlation is relatively low (Pearson correlation = 0.21). Figure 3a indicates that there is a large number of GPCR-endogenous ligand pairs where the evidence is strong (e.g. overall score > 0.5) for one but not the other partner, that is, evidence for disease involvement is often asymmetrically reported for one or other partner in these ligand-receptor pairs. It is possible that the involvement in the disease is not mediated through the partner interaction in such cases. However, since the identities of both partners in these interactions are well established, we should consider the evidence for the GPCR and its known endogenous ligand together as a pair to increase our confidence and supportive evidence for potential new target hypotheses. For example, genetic evidence for a disease association with an endogenous ligand may exist but the corresponding GPCR may turn out to be the better drug target due to, for example, druggability. To further quantify the added benefit of combining supportive evidence from GPCRs and ligands we first determined all disease-GPCR pairs and corresponding disease-ligand pairs with positive overall scores. If a pair had a positive score in only one category, we added the corresponding pair in the other category with 0 score. We then created joint disease-GPCR/ligand pairs and assigned a new overall score as the maximum of the scores from the disease-GPCR pairs and corresponding disease-ligand pairs. Figure 3b shows the distribution of the increase in overall score comparing the disease-GPCR pairs to the corresponding disease-GPCR/ligand pairs and Fig. 3c shows cumulative density function (CDF) for this change in score. While 93% of scores increased by 0.2 or less, many new high confidence pairs also emerged: 648 disease-GPCR/ligand pairs had a score of 0.5 or higher but did not have any supportive evidence (i.e. score = 0) for the corresponding disease-GPCR pair alone without considering the ligand. Of those, 355 pairs had a new score of 1.0 compared to the previous score of 0. Comparing the number of disease-GPCR/ligand pairs to the number of corresponding disease-GPCR pairs alone, the number of pairs without any evidence (i.e. score = 0) decreased by 62% and the number of high-confidence pairs (score > 0.5) was more than 1100 higher, a 69% increase (Fig. 3d).
GPCRs and endogenous peptidergic ligands and the highest stage in global pharma pipelines

Based on the past success of GPCRs as drug targets [2], GPCRs that have disease associations with a high score in the Open Targets Platform but are not currently pursued by the industry may potentially be high priority targets for the development of new therapies. Conversely, GPCRs with existing drug discovery programs as well as high scoring disease associations provide potential drug repurposing opportunities for compounds modulating these GPCRs if the top-ranked disease derived from Open Targets is different from the current indication pursued. To more closely examine this approach, we obtained a database of current drug discovery programs [24] and determined the highest stage in the drug discovery pipeline for each GPCR and endogenous peptide. Approximately half of the previously uniquely identified GPCRs and endogenous peptides had at least one program in the drug discovery pipeline (203 out of 397, and 209 out of 412, respectively; Fig. 4). We then stratified GPCR-disease pairs and endogenous peptidergic-ligand-disease pairs by the highest pipeline stage of the corresponding GPCR and peptide, respectively. Approximately 73% of GPCR-disease pairs with an overall Open Targets score below 0.5 had at least one program in the drug discovery pipeline for that target and this number increased to 93% for the GPCR-disease pairs with an overall Open Targets score of 0.5 or higher which corresponds to the 95th percentile of the overall score distribution as described above. In nearly 84% of such pairs, the GPCR has been recorded in at least one post-clinical stage of the drug discovery pipeline and only 6.5% of such pairs involve GPCRs without any drug discovery program (Fig. 4). Together, 56% of the GPCR-disease pairs involved GPCRs that had reached a clinical stage and those pairs involved GPCRs that had reached a clinical stage and those pairs also had significantly higher overall scores ($p = 3.8 \times 10^{-34}$, Wilcoxon rank sum test). Similarly, 47% of the ligand-disease pairs involved endogenous ligands that had reached a clinical stage and those pairs also had significantly higher overall scores ($p = 1.3 \times 10^{-29}$, Wilcoxon rank sum test). However, at least some of this relative over-representation of the late-stage pipeline among pairs with overall Open Targets score of 0.5 or higher may be driven by evidence resulting from the very same drug discovery programs as individual evidence types contribute differently to this enrichment. For example, we observed that genetic association and animal model evidence appears to be independent of pipeline status while literature evidence does not. For endogenous peptides,
the differences between lower Open Targets scores (< 0.5) and high Open Targets scores (≥0.5) are less prominent. For example, the endogenous peptide-disease pairs with an overall Open Targets score below 0.5 as well as the pairs with a score of 0.5 or higher both included approximately 29% of pairs where the ligand did not have any program in the drug discovery pipeline.

To illustrate how the Open Targets platform might be applied to prioritize a particular target-disease hypothesis for drug discovery, consider one of the examples listed in Table 2, GPR35 for inflammatory bowel diseases (IBD). The incidence and prevalence of IBD such as Crohn’s disease and ulcerative colitis are increasing over time globally [25] and estimates suggest that ~ 1.4 million people in the United States and 250,000 people in the United Kingdom suffer from this disease [25, 26]. The aetiology is currently not well known, and it is hypothesized that the genetically susceptible host suffers from compromised intestinal immune system response to commensal bacteria [25]. Currently, there is no known cure for this chronic condition, and constant care & symptomatic treatment is needed for patients suffering with this condition. Several genome-wide association studies have identified the GPR35 locus as one of the susceptibility loci for IBD [27, 28]. As the evidence listed in the Open Targets platform shows, GPR35 is currently investigated in clinical trials for pruritus and mastocytosis and presents a promising new therapeutic target for a number of disease indications including inflammatory and cardiovascular disease [29–33]. Currently, no drug targeting GPR35 is approved for IBD. Taken together, the evidence compiled in Open Targets strongly suggests that GPR35 could be investigated as a novel therapeutic option for IBD. It should be noted that lodoxamide, a GPR35 agonist, is an approved drug for conjunctivitis which could potentially be repositioned for IBD.

Another such example includes C C-X-C motif chemokine receptor 4 (CXCR4) as a potential new drug target for infectious diseases. The Open Targets platform identifies weak supporting evidence from RNA expression and genetic associations (score = 0.01 in each case) but strong pathway evidence. CXCR4 is part of the ‘Binding and entry of HIV virion’ pathway, a manually curated pathway from Reactome (score = 1.0), but is also listed in various relevant gene ontology pathways such as GO:0001618 ‘virus receptor activity’ which can easily be determined by following the link-out to the Uniprot database within the Open Target platform entry for CXCR4. The CXCR4 receptor is actually well known to play a critical role for the entry of the human immunodeficiency virus (HIV) into CD4+ T-cells but other viruses use this entry as well [34]. The literature text mining evidence shown in the Open Targets platform receives a score of 0.21. However, the platform identifies nearly 1200 publications further strengthening the hypothesis. Table 2 lists additional specific examples of possible repurposing opportunities and Additional file 1: Table S2 lists examples with overall score of 0.5 or higher.

### Discussion

Drug discovery and development programs focus on achieving therapeutic efficacy upon modulation of a specific drug

| Gene      | Disease term                  | Score |
|-----------|-------------------------------|-------|
| GPR35     | Inflammatory bowel disease    | 1.00  |
| CXCR4     | Infectious disease            | 1.00  |
| PTGER4    | Inflammatory bowel disease    | 0.90  |
| TSHR      | Graves disease                | 0.67  |
| NPSR1     | Asthma                        | 1.00  |
| LPAR6     | Alopecia                      | 1.00  |
| CELSR2    | Coronary heart disease        | 0.92  |
| GPR65     | Crohn’s disease               | 0.81  |
| TNFSF15   | Inflammatory bowel disease    | 1.00  |
| AGT       | Hypertension                  | 1.00  |
| IL17F     | Immune system disease         | 1.00  |
| CALCA     | Cardiovascular disease        | 1.00  |
| FBN1      | Vascular disease              | 1.00  |
| COL3A1    | Cardiovascular disease        | 1.00  |
| NPPA      | Cardiomyopathy                | 1.00  |
| IL33      | Respiratory system disease    | 0.77  |
target with a molecule in a patient population. A number of computational approaches to aid in target identification have been considered, and along with various bioinformatics resources they also point to the application of cheminformatics based approaches for ligand discovery [35]. However, the hypothesis that modulation of a specific target may potentially result in therapeutic benefit is often based upon years of scientific work which involves generating and/or accumulating experimental evidence in an iterative manner and then meaningfully integrating that information to further lend support to that hypothesis. The scientific evidence to build that hypothesis comes from multiple sources; and integration as well as evaluation of that data is critical for drug discovery programs. Especially in a world of rapidly growing data, the ability to integrate data from multiple sources with platforms such as Open Targets, presents an opportunity to systematically evaluate the available evidence to quickly generate hypotheses to identify targets that may further be followed up with additional experimentation [13]. It should be noted that the purpose of such efforts is not necessarily to identify novel target-disease associations per se but rather to prioritize such associations in order to identify the most promising opportunities for drug discovery.

In this study, we present the development and application of a systematic target identification approach on data from Open Targets platform. We first examine and characterize the distribution of the Open Targets score and its relationship with the individual evidence type scores. We then focus on a very successful target class of proteins in drug discovery, G-protein coupled receptors (GPCRs), along with their endogenous ligands. Specifically, we use a list of GPCRs and endogenous peptidergic ligands from IUPHAR, map them to Entrez gene identifiers, assemble data from various sources of evidence in the Open Targets platform and associate the disease terms with therapy areas for broader categorization. Although we are focusing on GPCR-ligand pairs in our analysis, our approach can be generalized to any heteromultimeric proteins or potentially to any pairs of proteins known to directly interact. Finally, we compare the Open Targets derived target-indication hypotheses (based on gene-disease associations) to the global pharmaceutical drug discovery landscape as a means to evaluate some of these hypotheses.

We observed that both GPCRs and endogenous ligands have a higher than expected number of associated disease terms in Open Targets. One explanation for this seemingly higher disease-relevance could be that these classes of proteins simply are better studied and understood than the proteome as a whole due to the extraordinary success of these protein classes as therapeutic drug targets [36]. The relatively high number of GPCR-disease pairs with an existing drug discovery program seems to confirm this view and suggests that this class of potential drug targets is well suited to evaluate the Open Targets score. We found that an Open Targets score of 0.5 corresponds to the 95th percentile of the overall score distribution and that over 90% of GPCR-disease pairs with a score of 0.5 or higher had a corresponding drug discovery program for that GPCR. This suggests that an overall Open Targets score of 0.5 could be used as a high confidence threshold when evaluating potential new target-indication hypotheses. We also found that confidence in such hypotheses was increased by more individual supporting evidence types.

Our current study highlights the benefit of combining different, independent sources of evidence supporting a target-disease hypothesis to increase confidence in its validity. This relationship was intentionally reflected in the design of the individual scores and, also in the overall score [13]. In particular, the overall score increases with the number of positive individual scores for a given target-disease hypothesis as shown in Fig. 2c. Another way the Open Targets platform can be used to accumulate existing supporting evidence is by combining data for closely related targets such as through a shared molecular pathway, heteromultimeric protein complexes, or through receptor-ligand pairs such as the examples highlighted in Table 1. It should be noted that the interpretation of an observed association between genes or proteins and diseases or medical conditions is not trivial. Such relationship may or may not be causal and it may be direct or indirect. Furthermore, it is often unclear if the disease association is due to an increase or decrease in activity or abundance of the functional protein. Some of the evidence integrated in the Open Targets database provides more clarity in this regard (e.g. availability of a known drug, Mendelian trait, knock-out animal model). In the context of GPCR-ligand relationships, it is also important to consider whether a ligand acts in a pathological or therapeutic role. For example, glucagon-like peptide-1 (GLP-1) can decrease blood sugar levels which has led to the development of GLP-1 receptor agonists as new drugs to treat type 2 diabetes [37]. Conversely, vasopressin plays a central role in the pathogenesis of hyponatremia which has led to the development of vasopressin receptor antagonists as a treatment [38]. As a result, each target-disease association of interest requires further careful evaluation of the evidence and subsequent experimental validation.

As with any systematic or global computational solution to a biological or biomedical problem, simplifications and generalizations are required. Therefore, a general approach applied to all disease terms and all potential drug targets such as the Open Targets platform may be more suitable in some situations than in others. For example, the current evidence presented by the Open Targets platform concentrates on data generated...
through methods that focus on the DNA or RNA level but
the action of a therapeutic drug is most often mediated at
the protein level, e.g. by disrupting protein-protein interac-
tions [39] or protein complexes comprised of multiple dif-
ferent genes [40]. In other cases, a protein might have
multiple splice forms, or both a membrane-bound and a
soluble form. In addition, in some cases the same gene
encodes multiple different peptides such as the GCG gene en-
coding glucagon, GLP-1, and GLP-2, each of which may
have different receptors as well as different disease associa-
tions. Additional evidence that reflects such complexities
could enhance the utility of the platform.

It should also be noted that as with any computational
approach, false positive and false negative results are un-
avoidable and should be expected. Each target-disease pair
merely represents a hypothesis that serves as a starting
point for drug discovery scientists looking to begin a new
research program. These hypotheses still require careful
evaluation, prioritization, and experimental validation.

Two final examples illustrate this point. Neuropeptide S
receptor 1 (NPSR1) was identified as a potential new drug
target for asthma in the Open Targets platform. Strong
genetic evidence supports the hypothesis [41–43] and the
Open Targets platform identifies over 60 publications sug-
gestng a role of NPSR1 in asthma but the exact mechan-
ism of NPSR1 in the disease remains elusive. Although
increased NPSR1 protein levels in plasma were reported
in asthma [44] and increased NPSR1 mRNA expression was
observed in eosinophils from severe asthmatic pa-
tients [45], experiments in an experimental asthma mouse
model showed no impact of Npsr1 deletion on airway in-
flammation or hyper-responsiveness, and the authors sug-
gested that NPSR1 affects the disease through a central
nervous system-mediated pathway [46]. Similarly, G
protein-coupled receptor 65 (GPR65), a receptor for psy-
chosine and several related glyosphingolipids, received a
strong Open Targets score for Crohn's disease mostly due
to its strong genetic association [28, 47, 48]. The protein's
role in the disease is not entirely clear but it may play role
in proton sensing [49] or acid sensing [50, 51] and may
regulate cytokine production of T cells and macrophages
[52, 53]. These examples further illustrate the importance
of systematically mining the Open Targets data and then
prioritizing target-indication pairs for follow-up experi-
mental work to validate the hypotheses.

Conclusion

In summary, by utilizing the Open Targets platform, data,
and evidence model, and by interrogation of underlying
and additional data, we have been able to generate various
GPCR – indication pair combinations, which form the basis
for development hypotheses for potential drug discovery
programs and this approach can be generalized in a
straightforward manner to include other drug target classes.

Methods

Open targets platform data

Open Targets gene-disease pairs and scores (September
9, 2017 version, Release 3.2; JSON format) were down-
loaded from the Open Targets website [13]. The data
download was parsed capturing disease term and Experi-
mental Factor Ontology (EFO) identifier, Ensembl gene
identifier and symbol, as well as 8 scores: overall, genetic
association, somatic mutation, known drug RNA expres-
sion, affected pathway, animal model, and literature min-
ning scores, respectively. Ensembl gene identifiers were
mapped to Entrez genes and official HUGO gene sym-
bols using relevant Bioconductor packages [54, 55].

Experimental factor ontology (EFO)

The EFO [56] was downloaded in OBO format (Septem-
ber 7, 2017 version). The ontology was parsed recursively
using the “is_a” relationships encoded in each entry in
order to determine one or more therapeutic areas for each
disease term. Specifically, an EFO term was considered a
therapeutic area if it was a directly associated with “dis-
ese” (EFO:0000408) through an “is_a” relationship. A
small number of such top-level terms were manually re-
mapped to a different therapeutic area (e.g. “heavy metal
poisoning”, “malignant epitheloid mesothelioma”, and
“sudden infant death syndrome”).

GPCRs and endogenous peptides

Three data tables were downloaded from IUPHAR
(http://www.guidetopharmacology.org) [19]: (a) a list of
GPCRs, (b) a list of endogenous peptides, and (c) the list
of all interaction data for endogenous ligands and their
GPCR targets. GPCRs were mapped to Entrez gene identi-
fiers by gene symbols, and endogenous peptides were
mapped to Entrez gene identifiers by Uniprot IDs using
relevant Bioconductor packages [54, 55] in both cases.

Comparison to Pharmapipeline

The Pharmapipeline database was retrieved from Informa
PLC [24]. It contains data on the current global pharma-
ceutical drug discovery pipeline and identifies drugs discov-
ery programs, their current status, molecular target, and
indication, among other data. Molecular target identifiers
were mapped to one or more Entrez gene IDs and drugs
without a matching gene identifier were removed from fur-
ther analyses. We summarized the drug discovery pipeline
stages as follows: none (“N/A”, global status 1), pre-clinical
(global status 2–5), clinical trial (global status 7–9), and
post-clinical (global status 10–13). For target-indication
pairs with multiple corresponding drug discovery programs,
we chose the highest stage as representative.
Additional file

Additional file 1: Table S1. Overall Open Targets score for disease-GPCR pairs and the corresponding disease-endogenous ligand pairs. Table S2. Possible repurposing opportunities with overall Open Targets score of 0.5 or higher. (XLSX 1930 kb)

Abbreviations

CDF: Cumulative density function; CXXC4: C-X-C chemokine receptor type 4; EFO: Experimental Factor Ontology; FDA: United States Food and Drug Administration; GLR2: Galanin receptor type 2; GCG: Glucagon; GLP-1: Glucagon-like peptide 1; GLP-2: Glucagon-like peptide 2; GPCR: G-protein-coupled receptor; GPR35: G-protein-coupled receptor 35; GPR65: G protein-coupled receptor 65; HIV: Human immunodeficiency virus; HUGO: Human Genome Organisation; IBD: Inflammatory bowel disease; NPSRI: Neuropeptide S receptor 1; RNAi: Ribonucleic acid

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Availability of data and materials

The datasets analysed during the current study are available in the Open Targets Platform, https://www.targetvalidation.org/downloads/data.

Authors’ contributions

JF, PS, ID, and DR conceived and designed the study. JF analysed and interpreted the data and wrote the manuscript. PS, ID, and DR critically revised and edited the manuscript. All authors have given final approval of the version to be published.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

JF, PS, and DR are employees and shareholders of GlaxoSmithKline. ID has no competing interests.

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