Data and text mining

Predicting cross-tissue hormone–gene relations using balanced word embeddings

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Abstract

Motivation: Inter-organ/inter-tissue communication is central to multi-cellular organisms including humans, and mapping inter-tissue interactions can advance system-level whole-body modeling efforts. Large volumes of biomedical literature have fostered studies that map within-tissue or tissue-agnostic interactions, but literature-mining studies that infer inter-tissue relations, such as between hormones and genes are solely missing.

Results: We present a first study to predict from biomedical literature the hormone–gene associations mediating inter-tissue signaling in the human body. Our BioEmbedS* models use neural network-based Biomedical word Embeddings with a Support Vector Machine classifier to predict if a hormone–gene pair is associated or not, and whether an associated gene is involved in the hormone’s production or response. Model training relies on our unified dataset Hormone-Gene version 1 of ground-truth associations between genes and endocrine hormones, which we compiled and carefully balanced in the embedded space to handle data disparities, such as between poorly- versus well-studied hormones. Our BioEmbedS model recapitulates known gene mediators of tissue–tissue signaling with 70.4% accuracy; predicts novel inter-tissue communication genes in humans, which are enriched for hormone-related disorders; and generalizes well to mouse, thereby holding promise for its extension to other multi-cellular organisms as well.

Availability and implementation: Freely available at https://cross-tissue-signaling.herokuapp.com are our model predictions & datasets; https://github.com/BIRDSgroup/BioEmbedS has all relevant code.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Inter-tissue communication forms the basis for life and health in multi-cellular organisms, and multiple organs/tissues are often affected in a systemic fashion in complex diseases [e.g. type 2 diabetes (Priest and Tontonoz, 2019) or cancer cachexia (Argilés et al., 2018)]. A grand goal of systems biology is to develop whole-body systemic models of healthy/disease conditions; and that calls for mapping not only the within-tissue biomolecular interactions, but also the crucial across-tissue interactions (Castillo-Armengol et al., 2019). Within-tissue interactions are focused heavily in current genomic or literature-mining studies—for instance, several recent methods that examine single-cell spatial omics data to predict ligand-receptor-based cell-cell interactions (Armingol et al., 2021; Türei et al., 2021) have focused largely on proximal inter-cell communication within a single tissue/organ, and not on inter-tissue interactions that are distal or endocrine in nature. Recently developed whole-body metabolic models for humans (Brul and Angione, 2019), such as the Harvey and Harvetta models [comprising 26+ organs and their metabolic interactions culled from literature and refined by genomic or other experimental data (Thiele et al., 2020)] are promising, but models of similar scale at the gene-level are lacking.

Emerging multi-tissue genomic datasets [e.g. the Genotype-Tissue Expression or GTEx (The GTEx Consortium, 2020) and Human Cell Atlas (Regev et al., 2017) data] have encouraged some gene-level efforts to address the dearth of studies on inter-tissue signaling, and there is a urgent need to augment/validate such
data-driven whole-body gene networks by literature-driven approaches. An ideal literature-mining system would extract relations of the form ‘Gene X in Tissue A interacts with Gene Y in Tissue B via a mediating signaling molecule’ from a literature corpus, and importantly also predict new relations using the semantic/syntactic attributes of the relevant genes and signaling molecules mentioned in the corpus.

In this work, we harness the large volume of biomedical literature [present in repositories like PubMed (Westergaard et al., 2018) with ~23 million abstracts] to identify genes involved in inter-tissue signaling mediated by hormones. We specifically focus on endocrine hormones, as they are a popular class of signaling molecules and endocrine biology has revealed and continues to reveal many hormones and their regulating genes. For example, in the pancreas tissue, gene INS produces the well-studied insulin hormone, which is processed and secreted into the blood with the help of other gene products and biomolecules; and the protein encoded by the INS gene is the primary receptor of the hormone in muscle and other tissues, where INS protein gets activated by the insulin hormone and affects several other genes downstream to regulate glucose uptake by body cells. A more recent example is a new type of vesicle-mediated communication between liver and adipose tissues involving non-protein-coding (microRNA) genes (Chen and Pfeifer, 2017; Zhao et al., 2020). The goal of this work is to extract all such hormone–gene relations, and classifying the hormone-associated genes as either source (genes aiding in hormone production/processing/secretion at a source tissue) or target (genes responding to the hormone directly via binding or as a downstream response at a target tissue).

Several challenges stand in the way of our goal of extracting hormone–gene relations from literature, including:

1. Lack of a unified database of ground-truth hormone–gene associations, since current hormone databases like HMBase (Rashid et al., 2009) and EndoNet (Donitz and Wingender, 2014) focus on the primary gene coding for a (peptide) hormone and the primary receptor genes, and not on the many other genes involved in hormonal processing/response.

2. Severe imbalance in known hormone–gene relations both in the number of source versus target genes of each hormone, and of associated genes per hormone, which varies widely across hormones (e.g. insulin is better studied than many other hormones).

3. Lack of standard in silico strategies for large-scale validation of novel hormone–gene predictions using independent data on inter-tissue signaling.

Our work attempts to address the barriers above to characterize inter-tissue signaling, and current studies have addressed only the second challenge above and that too in very different contexts/applications. Specifically, current literature-mining studies largely extracted relations among entities that are tissue-agnostic (i.e. does not depend on the tissue) or within-tissue (i.e. happens within one or more of the tissues or cell types), and includes relations, such as disease–gene (Bhasuran and Natarajan, 2018), gene–phenotype (Xing the tissues or cell types), and includes relations, such as disease–gene associations, since current hormone databases like HMBase (Rashid et al., 2009) and EndoNet (Donitz and Wingender, 2014) focus on the primary gene coding for a (peptide) hormone and the primary receptor genes, and not on the many other genes involved in hormonal processing/response.

We developed two classification models—BioEmbedS to predict hormone–gene associations, and BioEmbedS–TS to classify an associated gene into source versus target set of a hormone. These models, referred jointly as BioEmbedS*, are SVM classifiers trained and evaluated using our ground-truth dataset HGv1, and use word embeddings as input features (Fig. 1). An embedding of a word, such as a hormone name or gene symbol, is a vector representation of the word that is learnt via a deep learning model to capture the context of this word in a corpus. By using word embeddings, specifically the 200-dimensional vectors called BioWordVec (Zhang et al., 2019) derived from the PubMed corpus, we not only capitalize on this modern advance in literature mining, but also importantly balance the skew in our HGv1 dataset using established balancing techniques that can work only with feature vectors and not direct text (see Section 4 for details). The balancing step is crucial to prevent well-studied hormones with a large number of gene associations in HGv1 from unduly skewing our model predictions [Fig. 1 (inset histogram)].

Since a unified database of source and target genes for known hormones was not available, we expressly assembled such a database for 51 endocrine hormones, primarily ones listed in a Endocrine Society website (see Section 4), by integrating data from several sources (Donitz and Wingender, 2014; Rashid et al., 2009;
and Wingender, 2014; Rashid hormone–gene associations (Table 2) derived from the Appropriate GO term. We compiled a human dataset HGv1.human of about 2000 genes annotated to any species by taking the appropriate set of genes annotated to the term. We compiled a human dataset HGv1.human of about 2000 hormone–gene associations (Table 2) derived from the appropriate GO terms and augmented with primary genes (genes encoding a peptide hormone or hormone-binding receptors) from other sources (Donitz and Wingender, 2014; Rashid et al., 2009). Similarly, we constructed HGv1.mouse dataset by collecting mouse genes annotated to the GO terms collected in the first phase, and by taking the mouse homologs of the primary human genes. This work focuses on HGv1.human (simply referred to as HGv1 in the text), with HGv1.mouse dataset being used to study generalizability of our models.

2.2 Word embeddings are informative of relationship among hormones and genes

We first assess the quality of literature-based BioWordVec (Zhang et al., 2019) embeddings of hormone names and gene symbols using simple unsupervised learning methods. Hierarchical clustering of the word embeddings of the 51 hormones in our HGv1 dataset (Fig. 2a) revealed that functionally similar hormones often group together into clusters—for instance, neurotransmitter hormones like serotonin and dopamine are clustered together; and so are steroid hormones with sexual and reproductive functions, such as testosterone, estradiol and progesterone.

To evaluate the quality of embeddings of both hormones and genes, we predicted hormone–gene associations using the popular cosine similarity measure between word embeddings of a hormone and a gene, and obtained an average area under the receiver operating characteristics curve (ROC-AUC) of 0.69 (Supplementary Fig. S1). This result confirms the good quality of BioWordVec embeddings, seen in earlier studies on extracting relations among proteins and drugs (Zhang et al., 2019), in our new context of extracting hormone–gene relations from biomedical literature. This result also provides a baseline performance from a simple unsupervised method, against which we can compare the performance of our supervised BioEmbedS model. Furthermore, cosine similarity can aid interpretability of the hormone–gene pairs predicted by BioEmbedS—for instance, BioEmbedS-predicted genes GSTP1 and BRCA2 for the hormone estradiol, and Figure 2b shows how similar the embeddings of key estradiol receptor genes (ESR1 and ESR2) and estradiol-related disorders are to both the hormone and the predicted genes’ embeddings.

2.3 BioEmbedS strategy on disease–gene predictions is competitive with other methods

Due to lack of existing tools for hormone–gene predictions and due to several tools available for disease–gene predictions, we first validated our BioEmbedS strategy (of an SVM classifier trained on word embeddings) on predicting disease–gene associations from a

Table 1. Example snapshot of our HGv1 dataset: glucagon and adiponectin hormones along with their dominant source/target tissue(s) and GO terms

| Hormone source (HGF1) | Source terms | Target terms |
|-----------------------|--------------|--------------|
| Glucagon pancreas → liver | GO: 0070091 (glucagon secretion) | GO: 0033762 (response to glucagon) |
|                        | GO: 0004967 (glucagon receptor activity) | GO: 0031769 (glucagon receptor binding) |
| Adiponectin adipose → skeletal muscle, oocytes, pre-implantation embryos | GO: 0070162 (adiponectin secretion) | GO: 0033211 (adiponectin-activated signaling pathway) |

Note: ± denotes standard deviation here and elsewhere in the text.

Table 2. HGv1 dataset overview: summary of the 2009 hormone–gene associations in our HGv1.human dataset

| Number (#) of hormones/genes | 51/1453 |
|----------------------------|---------|
| Mean # of genes associated with a hormone | 39.39 (±78.37) |
| Mean # of hormones associated with a gene | 1.38 (±0.87) |
| # of source/target genes across all hormones | 519/1082 |
| Mean # of source genes for a hormone | 12.9 (±32.03) |
| Mean # of target genes for a hormone | 28.24 (±53.66) |

Note: ± denotes standard deviation here and elsewhere in the text.

Fig. 1. BioEmbedS model overview: our BioEmbedS model predicts if a hormone–gene pair is associated or not from D-dimensional word embedding vectors of the hormone name and the gene symbol. Our HGv1 dataset is crucial for systematic training/evaluation of our model, after its proper balancing to handle variability in available information for different hormones (see inset histogram; ‘assoc.’ stands for associated). In the toy-example shown, circles and triangles indicate hormone–gene pairs for two illustrative hormones; and below-the-boundary blue and above-the-boundary red symbols, respectively, denote the positive (associated) and negative (non-associated) genes for each hormone. The positive/negative classes are balanced across the two hormones, before separating them in a higher dimensional space using a SVM classifier. BioEmbedS-TS model has source and target genes for a hormone in place of positive and negative genes (A color version of this figure appears in the online version of this article.)
corpus called EU-ADR (Erik et al., 2012). For performance comparison, we used results reported by the BioBERT, BeFree and Joint Ensemble methods discussed before in Section 1; and our BioEmbedS approach is able to obtain a comparable F1-score of 85.84% relative to these methods (Table 3). This is promising as our approach, originally conceived for predicting hormone–gene links, performs reasonably well for a disease–gene prediction task. We do not delve further into these specific disease–gene predictions, as our main focus is to predict hormone–gene relations mediating inter-tissue signaling.

**Table 3. Performance on disease–gene predictions: our BioEmbedS approach’s 10-fold CV-based result is compared against existing methods’ reported results on the EU-ADR disease–gene corpus**

| Model                | Precision (%) | Recall (%) | F1-score (%) |
|----------------------|---------------|------------|--------------|
| BioEmbedS            | 77.13 ± 1.41  | 96.84 ± 2.67 | 85.84 ± 1.26 |
| BioBERT v1.0         | 81.05         | 93.90      | 86.51        |
| BioBERT v1.1         | 77.86         | 83.55      | 79.74        |
| BeFree               | 75.10         | 97.70      | 84.6         |
| Joint Ensemble learning | 76.43       | 98.01      | 85.34        |

Note: BioBERT results are obtained from Lee et al. (2020); results on BeFree and Joint Ensemble learning models are obtained from Bravo et al. (2015) and Bhasuran and Natarajan (2018), respectively. Bold-faced values indicate the best performance achieved in the corresponding column.

2.4 BioEmbedS predicts hormone–gene pairs reliably, and better than available alternatives

In our 5-fold cross-validation (CV) framework described in Methods, an SVM model with a third-degree polynomial kernel was chosen as a consistent classifier, and the resulting BioEmbedS models predicted hormone–gene associations with a reasonably good accuracy of 70.4% ± 1.8% and F1-score of 71.4% ± 2.7% (Supplementary Table S1). The SVM-based model also achieved better or comparable results than other classifier choices, such as logistic regression and decision trees (Supplementary Table S2). We also note that in our baseline comparison, our supervised BioEmbedS performed better than the unsupervised cosine similarity approach seen above (Supplementary Fig. S1).

Since there are no direct hormone–gene prediction tools available to which we can compare our method, the closest alternative was to match a (peptide) hormone to its primary gene encoding the hormone, and then use predicted associations of this gene to other genes. Predicted protein–protein and corresponding gene–gene associations are available in a widely used resource called STRING (Szklarczyk et al., 2015), and we found that our BioEmbedS’ hormone–gene scores were consistently better than STRING’s literature-mining-based scores for the corresponding (mapped) gene–gene pairs (Fig. 3a). We reiterate that STRING is not a direct hormone–gene prediction tool, rather it serves as a generic literature-mining baseline that is used to test if BioEmbedS can predict hormone–gene relations from literature relatively better. To
illustrate a bit more with insulin hormone as an example, whereas BioEmbedS would directly predict ‘Insulin–gene’ relations, STRING would predict ‘INS–gene’ associations where INS is the primary gene encoding the peptide hormone insulin, and using gene–gene scores from its ‘text mining’ evidence channel alone (i.e. ignoring STRING’s other evidence channels based on genomic context, experimental assays, etc., to allow STRING to be used as a literature-mining baseline; see Section 4 for more details).

We also evaluated our BioEmbedS model using word embeddings from the state-of-the-art BioBERT model (Lee et al., 2020) as input features. BioEmbedS using our default BioWordVec embeddings performed better than using the original BioBERT embeddings having 768 dimensions (downloaded from https://github.com/naver/biobert-pretrained) and the same embeddings reduced to 200 dimensions to match the dimension of BioWordVec embedding vectors (Supplementary Table S3).

### 2.5 BioEmbedS-TS classifies source versus target genes across different classes of hormones

For genes known to be associated with a hormone, we next classify if they are source or target genes for the hormone. The performance of our BioEmbedS-TS model was also reasonably good with accuracy of 79.0 ± 1.9%, and F1-score of 84.8 ± 1.3% for target genes and 66 ± 3.5% for source genes (see Table 4 and Fig. 3c). Superior performance in predicting target genes could be because of the disparity between the number of genes in the source and target sets. Table 2 shows that in source versus target classification, source genes constitute the minority class leaving us with fewer samples in the training data to learn underlying patterns.

To test if hormones with a large number of associated genes dominated our results, we grouped the hormones into bins based on the number of each hormone’s associated genes and calculated the bin-wise performance for BioEmbedS-TS as well as BioEmbedS. These results, reported in Supplementary Table S4 for BioEmbedS (with hormone-wise details in Supplementary File D1) and Supplementary Table S5 for BioEmbedS-TS, show that bin-wise performance is similar to the overall performance for most bins, thereby suggesting that our BioEmbedS* models perform well across a range of hormones.

### 2.6 BioEmbedS* models generalize reasonably to unseen hormones and well to another species

While previous results are already based on an unseen test set of hormone–gene links performed within a sound 5-fold CV framework, we also wanted to test how well our models would predict for fresh hormones that were never seen in the training/testing part of the CV framework. In other words, we tested BioEmbedS on an independent dataset of hormone–gene pairs pertaining to ‘unseen’ external hormones that were not used to train/evaluate the model (as these hormones had too few gene associations to be considered eligible in...
the final model building step and hence also every inner/outer CV iteration; see Materials and methods for details. This dataset contained 17 hormones forming 151 associated hormone–gene pairs and 148 non-associated hormone–gene pairs. Although, this dataset does not belong to the same distribution as the one used to train our model due to very few gene associations per hormone, BioEmbedS performed reasonably on this dataset obtaining accuracy of 65%, F1-score of 59% and ROC-AUC of 0.72 (Fig. 3b). Similarly, we applied BioEmbedS-TS on a set of hormone–source/target gene pairs from 40 unseen hormones, forming 501 hormone–target gene pairs and 188 hormone–source gene pairs. It was able to correctly classify these pairs with 69% accuracy and area under precision-recall curve (PR-AUC) of 0.43 (Fig. 3d). This PR-AUC of 0.43 is a drop from the average 0.722 seen in CV assessments in Table 4, however, it is higher than the random classifier’s PR-AUC of 0.27 (Fig. 3d), and hence in this regard, our model generalizes reasonably to unseen hormones with few known gene associations.

We also assessed how well our model trained on one species (human) generalizes to make predictions in another species (mouse). That is, we applied BioEmbedS model trained on hormone–gene pairs from the HGv1.human dataset (referred to as HGv1 so far) to predict mouse hormone–gene associations. Our BioEmbedS model was able to predict hormone–gene relations in the HGv1.mouse dataset at a reasonable accuracy of 71% and F1-score of 73%. Similarly, when BioEmbeds-TS model trained on HGv1.human was used to classify hormone-associated genes in mouse into source and target genes, we achieved 83% accuracy, and a F1-score of 78% for source genes and 87% for target genes. Since many human and mouse homologs have identical gene symbols (after converting to lower-case), we wanted to test if the good cross-species performance of our models could be driven solely by gene-symbols-based similarity of the HGv1.human and HGv1.mouse datasets. These results pertaining to hormones affecting a subset of tissues is shown in Figure 5. For instance, all the insulin predicted genes show that our BioEmbedS* models trained using human data generalize well to an organism other than human.

2.7 Novel gene predictions are enriched for the corresponding hormone-related diseases, and experimentally validated hormone-responsive genes

The promising performance of BioEmbedS seen so far encouraged us to apply BioEmbedS to predict association between each hormone in HGv1 and all 19 318 human protein-coding gene symbols (Braschi et al., 2019). We were able to predict many novel hormone–gene links not captured in HGv1, comprising new links to any of the 1453 genes in HGv1 (Table 2) or the remaining ‘out-of-HGv1’ genes, which were never seen during training/testing of our model. To validate these predictions, we tested whether the set of predicted genes for a hormone (at a default SVM probability score cutoff; see Section 4) was enriched for diseases already known [according to Pino et al. (2020)] to be related to the hormone. We found this was indeed the case for 16 of the 34 primary hormones (Supplementary File D2a, with primary indicating hormones in HGv1 considered eligible for training the final model as defined in Section 4), and 9 of the 17 unseen external hormones (Supplementary File D2b, with unseen referring to the remaining hormones in HGv1 with very few gene associations). These results pertaining to hormones affecting a subset of tissues is shown in Figure 5. For instance, all the insulin predicted genes are indeed significantly enriched for ‘Diabetes Mellitus’, a
Fig. 6. Disease enrichment in novel gene predictions: curves showing the number (no.) of known disease genes (as per DisGeNET; y-axis) recovered in top-k predicted genes (as per SVM score ranking; x-axis) of the corresponding hormone; focusing on all (lighter-shade red) versus novel (darker-shade black) predicted genes of the hormone. Our model (solid curves) performs better than chance recovery of disease genes by a random classifier (dashed lines). Only genes predicted for a hormone with SVM score>0 are considered here; all protein-coding genes are considered in Supplementary Figure S2 (A color version of this figure appears in the online version of this article.)

Table 5. Experimental validation of novel predictions: predicted (pred.) gene set refers to the set of novel (out-of-HGv1) target genes of a hormone predicted by our BioEmbedS* models (at default SVM probability cutoffs; see Section 4), and we compute its overlap with an experimentally derived reference (ref.) set of hormone-responsive differentially expressed genes (DEGs, including both up/down-regulated genes) already available for two well-studied hormones via gene expression omnibus (GEO) and Enrichr resources

| Hormone       | Ref. geneset name                  | Ref. experiment description               | Ref. geneset size | Pred. geneset size | Overlap size (overlap P-value) |
|---------------|------------------------------------|------------------------------------------|-------------------|--------------------|--------------------------------|
| Insulin       | Insulin Receptor Associates Promoters GSE107336 (Hancock et al., 2019) (GEO DEG FDR 5% via subseries GSE107334) | HepG2 cells treated with 10 nM insulin versus no insulin for 4 h | 4137              | 3424               | 1002 (2.74 x 10^-33)           |
| Insulin       | Insulin Receptor Associates Promoters GSE107336 (Hancock et al., 2019) (Enrichr geneset) | HepG2 cells treated with 10 nM insulin versus no insulin for 4 h | 457               | 3424               | 103 (0.00471)                  |
| Estrogen      | Estrogen human MCF-7 cells GSE11324 (Carroll et al., 2006) (Enrichr genesets ligand: 89,90,91) | MCF7 cells exposed to 100 nM estrogen for 3, 6 or 12 h versus 0 h | 953               | 9276               | 607 (1.40 x 10^-23)            |
| Estrogen      | Estradiol human estrogen receptor (ER)-positive MCF7 breast cancer cells GDS3217 (Len et al., 2007) (Enrichr genesets ligand: 39,40,41) | MCF7 cells exposed to 10 nM estradiol versus vehicle-only at 12, 24 and 48 h | 869               | 9276               | 487 (7.54 x 10^-7)             |

Note: GEO study identifiers are prefixed by ‘GSE’ or ‘GDS’. For insulin, besides the ref. geneset from Enrichr, the DEG table in GEO was used to derive another ref. geneset for the same experimental study at 5% false discovery rate (FDR). For estrogen, the DEGs in all timepoints of a study are combined to get a single ref. geneset per study.

disease term that is also recorded as insulin-related in the two hormone–ground-truth sources that we considered [Endocrine Society (https://www.hormone.org) and DisGeNET (Piñero et al., 2020) resources; see Supplementary D2a].

For well-studied hormones, such as insulin, we repeated the above analysis on only the novel predictions (i.e. predicted hormone–gene links not in HGv1) to test if the disease term enrichments were driven not just by known hormone-specific genes in HGv1 but also by the novel predicted genes. For insulin, BioEmbedS-predicted genes overlapped with 691 (of the 1507 Diabetes Mellitus genes recorded in DisGeNET (disease enrichment P = 9.55 x 10^-46), and 534 of these 691 overlapping disease genes were novel predictions (disease enrichment P = 4.85 x 10^-10). This trend of enrichment of predicted novel genes for the corresponding diseases can also be seen visually in Figure 6 across a range of cutoffs applied on the SVM score to call predictions—specifically, the curve for novel gene predictions is better than that of a random classifier for insulin and other hormones, and follows closely the overall gene curve for the most part. Furthermore, a more stringent subset of novel BioEmbedS predictions involving totally unseen out-of-HGv1 genes also got validated by a similar disease enrichment analysis (Supplementary Fig. S2).

Besides disease enrichment analysis, we can also test our novel hormone–gene predictions using a more direct evidence: experimentally validated ‘reference’ set of genes linked to a hormone. Experimental studies aimed at finding target rather than source genes of hormones are simpler to conceive, since a single transcriptomic (microarray or RNAseq) experimental study can reveal all the genes that are up/down-regulated in specific cell lines in response to treatment with a hormone. We could find such published experiments and associated hormone-responsive reference genesets for two of the four hormones in Figure 6 (see Section 4). For both these hormones, insulin and estrogen, the novel target gene predictions made by our BioEmbedS* models did indeed overlap with the tested experimentally derived reference genesets with high statistical significance (Table 5).

3 Discussion

This work elucidates the computational problems and challenges in the emerging area of inferring cross-tissue signaling interactions from biomedical literature, and presents a first approach BioEmbedS to specifically predict hormone–gene associations from biomedical literature with reasonably good absolute accuracy, and also comparable or better performance than other popular alternatives, such as STRING. Our BioEmbedS and BioEmbedS-TS models are enabled by a ground-truth dataset HGv1 that we carefully compiled and balanced across different stratifications of the training data in the space of mapped word embeddings of hormone names and gene symbols. The better performance of our method over other alternatives follows from our models being the first systematically developed ones to make hormone–gene predictions, and other methods being literature-mining-based relation extraction methods that are generic or designed for other prediction tasks, such as disease-gene prediction.
Our HGv1 dataset can be viewed as a two-layered/bipartite graph (with HGv1 hormone–gene relations being the edges/links between nodes in the hormone layer and the gene layer), hence our hormone–gene prediction task can be viewed as a link prediction problem in bipartite graphs (Kunegis et al., 2010). Current link prediction methods for bipartite or more general graphs (Kumar et al., 2020; Lu and Zhou, 2011), such as ones based on pairwise node similarity, utilize attribute data available at the nodes and/or the structural connectivity/neighborhood of nodes based on existing links in the graph to predict new links. We decided to utilize only the attribute data (word embeddings) of nodes to do hormone–gene bipartite link prediction, since our HGv1 graph does not have rich structural connectivity. In detail, a large number of genes in our HGv1 dataset (1102 of the 1453) are uni-hormone genes connected to only one hormone, and removing the few other multi-hormone genes disconnects the HGv1 bipartite graph into several connected components (one per hormone).

A caveat in this work worth mentioning is that randomly selected genes for a hormone need not be truly negative examples. Our model may also predict false associations based on high co-occurrence or context word similarity but no functional relationship, and a careful set of negative examples can help mitigate this issue. Our balancing strategy using Synthetic Minority Oversampling Technique (SMOTE) is also not without its pitfalls, especially when applied to high-dimensional data (Fernández et al., 2018), and we coupled it with under-sampling to carefully balance our training data across different hormones that are represented in the literature to different extents and to address imbalance in the number of source versus target genes. Also, we use relatively well-studied ‘primary’ hormones (with a higher number of known hormone-gene associations) for training the model. So when validating the method on independent test set comprising poorly studied hormones with fewer known hormone–gene associations, there is an expected drop in performance. Nevertheless, reasonably good accuracy of our model on different unseen test sets and on an organism other than human, and our disease enrichment and experimental validation analysis of novel gene predictions taken together suggest that our model predictions are indeed generalizable to make novel predictions about inter-tissue gene signaling mediated by a hormone.

Future work could focus on integrating literature information with independent omics data, such as multi-tissue genomic data (The GTEx Consortium, 2020) or rapidly accumulating single-cell/spatial transcriptomics data (Armingol et al., 2021). This literature-omics integration could not only improve model performance, but also help interpret further our novel hormone–gene predictions. For instance, a location (tissue or cell type) where a hormone–gene association happens could be predicted using the corresponding location’s omics data. As another example, systematic interpretation/validation of non-coding gene predictions for hormones [such as the preliminary long non-coding RNA (lncRNA) predictions, we provide in our website] using independent omics data could potentially reveal unexplored routes of inter-tissue communication. Finally, a special feature of our ground-truth hormone–gene dataset HGv1 is its applicability to organisms beyond human and mouse as it is based on species-agnostic GO terms, and this bodes well for extending our work to other multi-cellular organisms in the future. We hope our work on inter-organ hormone–gene network models, demonstrated concretely for the human and mouse organisms, stimulates more work along these lines on cross-tissue signaling for different organisms and thereby advances the field of whole-body, cross-tissue gene network modeling.

4 Materials and methods

4.1 Our BioEmbedS and BioEmbedS-TS approach

Word-embeddings-based classifiers: To proceed with the classification models, we need to represent our data (genes and hormones) in a way that can be used by a classifier. There are several ways of doing it, the most recent and efficient methods being from the deep learning community, known as word embeddings. Specifically, we use word embeddings for gene symbols and hormones from a FastText model pre-trained on the PubMed biomedical corpus called BioWordVec (Zhang et al., 2019) (BioWordVec model/ embeddings are downloaded from https://github.com/ncbi-nlp/BioSentVec). FastText (Bojanowski et al., 2017) is a neural network model learned by minimizing cross-entropy loss between each word and its predicted context within a fixed window size. We used 200D embeddings obtained using a skip-gram-based implementation with window size 20. This ensures the embedding vectors of not only co-occurring entities in a document but also entities with similar word neighborhoods exhibit high similarity (Park et al., 2019). BioWordVec actually uses subword information to obtain embeddings of words (including out-of-vocabulary words), i.e. the embedding of each word is represented by the sum of embeddings of all n-grams (3 \leq n \leq 6) in the word (after converting the words, including hormones and gene symbols, to lower-case) (Bojanowski et al., 2017; Zhang et al., 2019). Please note that BioWordVec is already trained using the PubMed corpus and so our HGv1 dataset is not used to obtain the word embeddings; HGv1 (training/testing splits) is instead used along with the pre-trained word embeddings to build our BioEmbeds* classifiers.

Existing studies on prediction of disease–gene associations have shown that SVM classifiers perform well when used with word embeddings (Bhasaran and Natarajan, 2018). So we explored SVM along with Random Forest (RF) classifiers as our primary classifiers, and compared them with other secondary choices of classifiers as well, and found SVMs to perform well (see Section 2). Hence, we decided on an SVM-based model to predict hormone–gene relations and call it BioEmbedS (with S denoting SVM). Similarly, we use an SVM-based model to classify source versus target genes for associated genes for a given hormone and call it BioEmbedS-DS (with -DS denoting Target versus Source).

Stratified/nested CV: We explored the parameter space (both hyper-parameter tuning and parameter learning) of our SVM and RF models using a 5-fold CV strategy that is stratified to ensure even distribution of each hormone’s genes across the different folds and nested to allow proper partitioning of the HGv1 data into training, validation and test sets. In detail, a stratified split amounts to considering each hormone with a certain number of associated genes in HGv1 (denoted n), and putting randomly chosen \( \frac{n}{k} \) genes into each of the 5-folds, and the remainder genes randomly (one each) into any of the 5-folds. This procedure done for the BioEmbedS model ensures that each fold has genes belonging to each hormone proportional to their presence in the overall dataset. A similar procedure to distribute the number of source (and separately target) genes of each hormone evenly across the 5-folds was done for the BioEmbedS-TS model as well. For BioEmbedS, we also created a negative class (non-associations) dataset of the same size across hormones and folds as the positive class (hormone–gene associations) dataset described above. In detail, for each hormone, we excluded its set of positive/associated genes (of size denoted by \( n \)) from all genes present in HGv1, and randomly chose from the remaining genes a set of \( n \) ‘negative’ genes, which are then split across the five CV folds in the same manner as the positive genes for this hormone. In the nested 5-fold CV strategy that we use to build both BioEmbedS and BioEmbedS-TS models, we make a train/validation/test split using 3/1/1-folds, respectively, and identify the classifier’s parameters by using only train and validation splits. The test split is kept aside from the training process, and used solely to report the model performance.

Over/under-sampling (balancing) training set embeddings: Our HGv1 dataset has a skewed distribution of the number of genes associated with different hormones [Fig. 1 (inset histogram); Table 2], so it is important for our model to prevent well-studied hormones from having a large number of gene associations in HGv1 from unduly influencing our model. To address the problem of class imbalance, oversampling techniques like SMOTE (Chawla et al., 2002) synthesize new examples from existing ones for the minority class, whereas under-sampling techniques like Condensed Nearest Neighbours (Hart, 1968), TOMEK Links (Tomek, 1976), etc. selectively remove examples from the majority class. A combination of oversampling and
under-sampling techniques is shown to perform better than using either alone (Chawla et al., 2002). We strategically apply a combination of SMOTE and TOMEK Links on the mapped embeddings of only genes with exactly one hormone association in HGv1. Working in the space of embeddings of such ‘uni-hormone’ genes is both desirable in facilitating SMOTE oversampling of our data, and permissible as a large fraction of all genes in HGv1, 1102 of all 1433, are uni-hormone (i.e. associated uniquely with some hormone, as opposed to being multi-hormone or associated with more than one hormone within HGv1). This technique applied to BioEmbedS and a similar technique applied to BioEmbedS-TS handle the large variation in HGv1 in the number and type (source versus target) of genes respectively across hormones.

For BioEmbedS, we assign a unique class ID to every hormone, and all the genes uniquely associated with the hormone belong to the class indicated by its class ID (with all multi-hormone genes discarded). We then use a combination of SMOTE oversampling and TOMEK Links under-sampling [with the number \( k \) of nearest neighbors in SMOTE set to two in the implementation used (Lemaitre et al., 2017)] to get an approximately equal number of genes/examples (same as the highest number of genes associated with a hormone among all the hormones in HGv1) for every hormone. We now have a dataset with approximately equal numbers of the genes related to every hormone that forms hormone–gene pairs of positive class for our binary classification problem. The following strategy is applied to create the negative class (set of non-associated hormone–gene pairs). For every hormone, we construct a set that contains the ‘gene–hormone’ (source/target) examples found in SMOTE (oversampling and under-sampling) associated with all the hormones except the one under consideration. We randomly select from this negative set as many examples as in the positive set of this hormone, in order to maintain class balance between the positive and negative associations for the hormone. Repeating this process for each hormone finally results in a dataset that is balanced across the different hormones and positive-negative class.

For BioEmbedS-TS, we define two unique IDs for every hormone; one maps to the source genes and the other maps to the target genes for that hormone (again all multi-hormone genes, defined exactly the same way as above for BioEmbedS, are discarded, along with the small set of genes that are annotated as both source and target for the same hormone). The source and target genes associated with a hormone belong to classes indicated by their IDs. This makes the number of classes equal to twice the number of hormones present in our dataset. We use SMOTE and TOMEK Links strategy as for BioEmbedS to get an approximately equal number of source and target gene examples (as the highest number of number of source/target genes associated to a hormone among all the hormones in HGv1) for every hormone. Finally, we have a dataset that is balanced across the different hormones and source/target class.

Putting it all together—balancing and model selection within nested CV: The overall careful application of our balancing and model selection steps to the 3/1/1 train/validation/test folds is shown in Algorithm 1, along with what constitutes the training and testing sets of each inner/outer CV iteration (e.g. validation and test folds are the respective testing sets for inner and outer CV loops). As for the genes considered by Algorithm 1, training sets are restricted to contain only uni-hormone genes to facilitate application of SMOTE and TOMEK Links as discussed before, whereas testing sets are allowed to contain both uni- and multi- hormone genes to reflect a real-world setting where any gene may be queried for its association to a hormone. As for the genes considered for the BioEmbedS model, the overall algorithm considers only hormones with at least five gene associations to permit 5-fold CV; additionally, each inner/ outer CV iteration considers only hormones with sufficient gene associations as ‘eligible’ for further analysis. In detail, every hormone with at least three associations to uni-hormone genes in an iteration’s training set is considered eligible for this iteration (to permit two-nearest-neighbor SMOTE), and the remaining hormones are removed from this iteration’s training as well as testing sets. Similarly for the BioEmbedS-TS model, hormones with at least five source and five target gene associations are considered for 5-fold CV. In each inner/outer CV iteration’s training set, only the hormones with at least three uni-hormone–source gene and three uni-hormone–target gene associations are considered ‘eligible’, and the remaining hormones are removed from training as well as the test set for this iteration.

SVM and RF models with different hyper-parameter combinations are trained on the resulting balanced training folds, and Cohen’s Kappa score of these trained models on the validation folds is used to select the best model. In each iteration, the test fold is never used for training or choosing hyper-parameters for the classifier. For BioEmbedS, an SVM model with a third-degree polynomial kernel was chosen as the best classifier by Algorithm 1 (Step 10) consistently in all five outer CV iterations (Supplementary Table S1). This SVM model was later trained using the entire HGv1 dataset, after applying similar restrictions and balancing as in Step 11 of Algorithm 1, to obtain the final BioEmbedS model—that final model was used for making all protein-coding gene predictions (including novel ones discussed in Section 2). Hormones found eligible for building the final BioEmbedS model amounted to 34 and are known as primary hormones hereafter; the remaining 17 ineligible hormones would also have been ineligible (and hence discarded or unseen when choosing/training models) in each of the inner/outer CV iterations of Algorithm 1 due to these hormones’ insufficient gene associations. For BioEmbedS-TS, SVM classifiers with different kernels/hyper-parameters were chosen as the best classifier in the different outer CV iterations of Algorithm 1 (Step 10); Among these best classifiers, we chose the simplest model: an SVM classifier with a third-degree polynomial kernel. We then trained this SVM model using the entire HGv1 dataset of source versus target hormone–gene relations, after applying similar restrictions and balancing as in Step 11 of Algorithm 1, to obtain the final BioEmbedS-TS model—that final model was applied to classify all hormone–gene associations predicted by BioEmbedS as source versus target type (using cutoffs shown below at the end of Section 4). Only 14 hormones with sufficient source and target gene associations were found to be eligible for building the final BioEmbedS-TS model.

4.2 Evaluation framework: performance metrics and enrichment/clustering analysis

We evaluate our model on five different unseen test sets (as shown in Algorithm 1), by reporting the performance of classifiers on these test sets using standard performance metrics like precision, recall, F1-score, accuracy, Kappa score, ROC-AUC and PR-AUC. All these metrics take highest value in the range \([0,1]\) and hence can be expressed as percentages (with the exception of Kappa score that can take negative values), with higher values indicating better performance. Supplementary Methods S1.1 provides definitions for these metrics assuming the two classes in our binary classification problems are as follows: for BioEmbedS, we naturally let positive and negative class to refer respectively to association and non-association of a hormone–gene pair; for BioEmbedS-TS that starts with an associated hormone–gene pair, we arbitrarily let positive and negative class refer to hormone–source gene pair and hormone–target gene pair association, respectively. Given these class definitions, a false positive for instance, for BioEmbedS would be a non-associated hormone–gene pair that is wrongly predicted as associated by our method; and for BioEmbedS-TS would be a hormone–target gene pair that is wrongly predicted as hormone–source gene pair by our method.

Hormone–gene predictions are called using the final BioEmbedS model described above at an SVM probability score of at least 0.7, with this default cutoff value chosen so as to get a reasonable number of predictions for each hormone. To validate the predicted genes of a hormone, we perform a disease enrichment analysis using the Enrichr tool (Chen et al., 2013) and DisGeNET (Piñeiro et al., 2020) collection of known disease-related genes. All reported disease enrichment \(P\)-values from this analysis are corrected for multiple testing of different DisGeNET disease terms. For validation of novel predictions, besides enrichment for disease genestes, we also tested for overlap with experimentally derived reference genestes. We
obtained these reference sets by searching the names of each of the four hormones in Figure 6 against all terms in the following Enrichr libraries of reference gene sets (Chen et al., 2013): ‘Ligand Perturbations from GEO’ (‘down’ and ‘up’ gene sets combined, since, we do not make distinction between down/up-regulated target genes in our predictions), and ‘RNAseq Automatic GEO Signatures Human’ (again ‘down’ and ‘up’ gene sets combined) (https://maayanlab.cloud/Enrichr/#libraries, accessed Jun 30, 2022). We excluded studies where the hormone was used in conjunction with some other stimuli to treat the cells, and we also chose cell line over disease studies (e.g. for insulin, we preferred HepG2 cell line hormone-treatment over insulinoma-tumor-related studies). These choices helped avoid any unwanted influence from non-hormonal stimuli or disease factors when identifying a hormone’s target genes. Hypergeometric distribution is used to obtain the Overlap P-values reported in Table 5.

Also, as part of validating our predictions, we inspected the pairwise overlap between the highly significant gene sets predicted for hormones, and found that biologically related hormones, such as insulin, leptin and glucagon, indeed group together in a hierarchical clustering-based dendrogram (Supplementary Fig. S3), due to their higher pairwise similarities than most other hormone pairs.

In comparison of STRING (Szklarczyk et al., 2015) to our BioEmbedS model within the 5-fold CV framework, STRING is used as a generic literature-mining baseline as it is not a direct hormone–gene prediction tool. We evaluated STRING on the following 14 peptide hormones alone (primary gene encoding the hormone shared in parenthesis): calcitonin (CALCA), cholecystokinin (CCK), glucagon (GCG), insulin (INS), leptin (LEP), parathyroid hormone/parathyrin (PTH), thyrotropin-releasing hormone (TRH), antidiuretic hormone/vasopressin (AVP), prolactin (PRL), somatostatin (SST), gastrin (GAST), ghrelin (GHRL), growth hormone-releasing hormone (GHRHR) and oxytocin (OXT). A ranked list of STRING’s hormone–gene predictions were obtained using the text-mining scores (in the 9606.protein.links.detailed.v11.0.txt file downloaded from the STRING website) for genes associated by STRING to each hormone’s primary gene (and assuming the lowest score of zero for non-associated genes). The same five CV folds derived from the HGv1 dataset were used to evaluate both BioEmbedS and STRING, after excluding positive/negative examples involving (i) hormones that were not eligible in the fold (see Algorithm 1) when evaluating both BioEmbedS and STRING, and (ii) hormones not present in the above peptide hormone list when evaluating STRING alone.

4.3 Implementation details and data/code availability

Implementations are done using Python Scikit-learn framework using decision_function() and predict_proba() methods of support vector classification in sklearn respectively to obtain the SVM model scores for all hormone–gene pairs and the probability of association between hormone–gene pairs (probability score of SVM). For reproducibility purposes, we provide hyper-parameter choices in Supplementary Methods S1.2 and open-source code and relevant data in a public repository. Specifically, the proposed HGv1 dataset along with our models’ predictions, and the code to reproduce this work are available respectively at the web portal https://cross-tissue-signaling.herokuapp.com/, and code repository https://github.com/BIRDSgroup/BioEmbedS. In the web portal, we show BioEmbedS-predicted hormone–gene associations at the default cutoff (of at least 0.7 for BioEmbedS SVM probability), as well as BioEmbedS-TS-predicted type of the association (i.e. source versus target type for hormones that were used to train the BioEmbedS-TS model; as explained above, hormones with insufficient number of source or target genes were not used to train the BioEmbedS-TS model). In detail, a hormone-associated gene whose BioEmbedS-TS SVM probability is: (i) at least 0.7 is predicted as a source gene for the hormone; (ii) at most 0.3 is predicted as a target gene for the hormone; and (iii) the rest are called as inconclusive or ‘don’t know’ genes. This association type of source or target can then be used to select either the known source or target tissues respectively of the hormone, as recorded in HGv1, to infer the potential locations (tissues) of a hormone–gene association. Our web portal also lists these predicted tissue locations. Furthermore, the target genes obtained using the above default cutoffs were used toward the experimental validation analysis reported in Table 5.

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Author contributions

A.J. and M.N. formulated the study and overall modeling approaches; A.J. implemented the training/testing of all primary models and performed associated analyses; T.K. and M.N. compiled the HGv1 human dataset, and provided inputs on the modeling approaches; T.K. assembled the HGv1 mouse dataset, and studied novel gene predictions at the hormone and inter-tissue level; M.R. performed associated analyses of secondary modeling approaches, visualized embeddings, and developed the website with inputs from A.J. and other co-authors; T.L. performed disease and pathway enrichment, hierarchical clustering and exploratory bioRNA analyses; A.J., M.N., T.K., T.L. and M.R. interpreted results; A.J., M.N. and T.K. wrote the manuscript; M.N. guided and supervised the study.

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