

Evaluating Methods for the Prediction of Cell Type-Specific Enhancers in the Mammalian Cortex

Authors:

Nelson J. Johansen^{1,11}, Niklas Kempynck^{2,11}, Nathan R. Zemke³, Saroja Somasundaram¹, Seppe De Winter², Marcus Hooper¹, Deepanjali Dwivedi¹, Ruchi Lohia⁴, Fabien Wehbe⁵, Bocheng Li⁶, Darina Abaffyová², Ethan J. Armand⁷, Julie De Man², Eren Can Eksi², Nikolai Hecker², Gert Hulselmans², Vasilis Konstantakos², David Mauduit², John K. Mich¹, Gabriele Partel², Tanya L. Daigle¹, Boaz P. Levi¹, Kai Zhang^{6,8}, Yoshiaki Tanaka⁵, Jesse Gillis⁴, Jonathan T. Ting^{1,9}, Yoav Ben-Simon¹, Jeremy Miller¹, Joseph R. Ecker¹⁰, Bing Ren³, Stein Aerts², Ed S. Lein¹, Bosiljka Tasic¹, Trygve E. Bakken^{1,12}

¹Allen Institute for Brain Science, Seattle, WA 98109.

²VIB Center for AI & Computational Biology, VIB-KU Leuven Center for Brain and Disease Research & KU Leuven Department of Human Genetics, Leuven, Belgium.

³Center for Epigenomics, Department of Cellular and Molecular Medicine, University of California, San Diego, La Jolla, CA 92093.

⁴Physiology Department and Donnelly Centre for Cellular and Biomolecular Research, University of Toronto, Toronto, Ontario, Canada.

⁵Maisonneuve-Rosemont Hospital Research Centre, University of Montreal, Montreal, Quebec, Canada.

⁶School of Life Sciences, Westlake University, Hangzhou, Zhejiang, China.

⁷Bioinformatics and Systems Biology Program, University of California, San Diego, La Jolla, CA 92093.

⁸Westlake Laboratory of Life Sciences and Biomedicine, Hangzhou, Zhejiang, China

⁹Department of Physiology and Biophysics, University of Washington, Seattle, WA 98195.

¹⁰Salk Institute for Biological Studies, La Jolla, CA 92037.

¹¹These authors contributed equally

¹²Lead contact

Correspondence: trygveb@alleninstitute.org

Summary:

Identifying cell type-specific enhancers in the brain is critical to building genetic tools for investigating the mammalian brain. Computational methods for functional enhancer prediction have been proposed and validated in the fruit fly and not yet the mammalian brain. We organized the ‘Brain Initiative Cell Census Network (BICCN) Challenge: Predicting Functional Cell Type-Specific Enhancers from Cross-Species Multi-Omics’ to assess machine learning and feature-based methods designed to nominate enhancer DNA sequences to target cell types in the mouse cortex. Methods were evaluated based on *in vivo* validation data from hundreds of cortical cell type-specific enhancers that were previously packaged into individual AAV vectors and retro-orbitally injected into mice. We find that open chromatin was a key predictor of functional enhancers, and sequence models improved prediction of non-functional enhancers that can be deprioritized as opposed to pursued for *in vivo* testing. Sequence models also identified cell type-specific transcription factor codes that can guide designs of *in silico* enhancers. This community challenge establishes a benchmark for enhancer prioritization algorithms and reveals computational approaches and molecular information that are crucial for the identification of functional enhancers for mammalian cortical cell types. The results of this challenge bring us closer to understanding the complex gene regulatory landscape of the mammalian brain and help us design more efficient genetic tools and potential gene therapies for human neurological diseases.

Keywords:

AAV, enhancer, multiome, RNA-Seq, ATAC-seq, DNA methylation, HiC, mouse, primates, cortex, cell types, machine learning, benchmark, challenge

Introduction

The mammalian neocortex, responsible for higher-order cognitive functions and sensorimotor processing, includes the primary motor cortex (M1), which facilitates fine motor movement and is composed of diverse cell types with distinct molecular signatures^{1,2}. Some neurodegenerative diseases, including Parkinson's, Huntington's, and amyotrophic lateral sclerosis (ALS), affect specific M1 cell types and result in impaired coordination and dexterity³. There is an urgent need for genetic tools to selectively access vulnerable cell populations to probe cortical circuit function and treat disease. To this end, cell type-selective candidate enhancers have been identified based on single cell genomic profiling of mouse cortex and have been used to create recombinant adeno-associated virus (AAV) vectors to drive exogenous transgene expression in the predicted target cell types⁴⁻⁶. With increasingly comprehensive molecular phenotyping of the brain⁷⁻¹¹, there is the prospect of developing AAV tools to target a wide diversity of cell types across the brain.

Identification of cell type-specific viral tools remains challenging because experimental validation is low-throughput and expensive. A recent study by Ben-Simon et al. 2024⁶ tested 682 enhancers selected based on specificity of open chromatin within a cortical cell type of interest and achieved an overall average success rate of 30%. Consequently, the field needs new computational approaches to improve functional enhancer prediction⁹ and accelerate viral tool development that integrates comprehensive molecular profiling within and across species. However, it remains poorly understood how well different approaches predict functional over non-functional enhancer genomic sequences. Community-guided challenges have demonstrated their effectiveness in rigorously evaluating novel computational methods and advancing knowledge in the genomics field¹². To date, there has been no community-led effort to address the prioritization of cell type-specific enhancers using cutting edge cell type-resolved atlases from multiple species.

In this study, we present the BICCN Challenge, where six teams from computational biology labs across the world participated to predict functional cortical cell type-specific enhancers. We introduce a community-driven benchmark and metrics to identify top-performing approaches that prioritize functional, cell type-specific viral tools. By investigating the methodologies and biological priors used by high-performance methods, we aim to contribute to the refinement of functional enhancer prediction to selectively target cell types in the mammalian brain.

Results

A community challenge to predict functional enhancers

We provided teams with a comparative multi-omics study of M1 by Zemke et al.¹³ that measured the molecular profiles of individual nuclei using single-cell multi-omics and single-cell methyl-Hi-C (snm3C) in human, macaque, marmoset and mouse (**Fig. 1A**). Multiple species were included because molecular and DNA sequence patterns associated with enhancer activity are conserved in the mammalian brain^{1,14}. Teams were asked to combine cross-species genomics measurements and biological priors to prioritize cell type-specific and functional enhancer elements (**Fig. 1B**).

Teams' predictions were evaluated against 677 recombinant adeno-associated virus (AAV) vectors among 682 vectors that were designed to label 19 M1 cell subclasses and assessed for *in vivo* enhancer specificity and brightness in the mouse brain⁶. Validation data for five enhancers were released after the challenge and were therefore excluded from the analysis. The cell type-specificity in the neocortex of each validated enhancer virus was assessed through manual evaluation of epifluorescence images, following the methodology outlined in Ben-Simon et al.⁶. The enhancers were classified into four groups: (1) On-Target (N=202), specific labeling of the targeted cell type; (2) Off-Target (N=96), labeling of non-targeted cell types in the neocortex; (3) Mixed-Target (N=100), labeling of targeted and non-targeted cell type(s); and (4) No-Labeling (N=279), no fluorescence in the mouse neocortex (**Fig. 1C, Supplemental Table 1**). To further quantify the *in vivo* activity of 191 enhancers, SYFP2-positive cells were extracted from the primary visual cortex (V1) and analyzed with Smart-seq v4 (SSv4) sequencing⁶.

The challenge was run over a 3-month period. Participants submitted enhancer lists at several intervals, and performance was reported on a public leaderboard¹⁵. In the final evaluation round, teams provided a detailed description of their approach (**Methods**). To rigorously evaluate methods, we developed an evaluation metric that is optimized when enhancers with On-Target activity are ranked highest, and Mixed-Target, Off-Target and No-Labeling enhancers are ranked lowest for the respective cell type (**Fig. 1D, Methods**). Final scores per method were computed based on both epifluorescence and SSv4 metrics using the entire corpus of validated enhancers. We are hosting both the evaluation metric and an automated scoring tool via GitHub¹⁵ for the community to fairly evaluate novel enhancer prioritization methods.

Top performing submissions focused on ATAC-seq specificity

Challenge participants from 5 teams contributed 79 submissions that comprised 16 unique enhancer prioritization methods, and the ArchR¹⁶ method was included as a performance baseline. We grouped the methods into five broad categories based on the included enhancer features: (1) ATAC-seq (meta), (2) Enhancer codes, (3) Feature Ranking, (4) Sequence Model and (5) Integration Model (**Fig. 1E**). The top three teams (Aerts, ArchR and PeakRankR) achieved comparable performance with normalized scores between 0.36 and 0.41, while the remaining teams achieved scores below 0.27 (**Fig. 1F**). Teams submitted diverse approaches and markedly improved during the challenge (**Fig. 1E-F, Supplemental Fig. 1, Supplemental Table 2**).

High-performing teams used similar approaches that leveraged ATAC-seq features, including differential chromatin accessibility (specificity) and signal strength at a given enhancer genomic location. The top performing submission gained a slight advantage by including RNA-seq and leveraging SCENIC+¹⁷ to predict cell type-specific transcription factor (TF)-enhancer-gene triplets. The runner-up baseline method, ArchR, used careful selection of background cells to minimize biases such as transcription start site (TSS) enrichment when performing pairwise statistical tests. The third-ranking team, PeakRankR, calculated three ATAC-seq metrics – specificity, magnitude and coverage – that were combined to discriminate cell type-selective enhancers.

Comparison of team submissions highlighted the genomics data, biological priors, and methodology that enabled accurate prediction of functional cell type enhancers (**Supplemental Fig. 2**). While all submissions used mouse ATAC-seq data, they varied in how the signal was normalized to handle batch effects and how cell type specificity was calculated. Surprisingly, inclusion of DNA-methylation, chromatin folding (HiC), or primate data decreased performance, potentially due to increased model complexity and overfitting. Also, the Tanaka team used all data types but retained only the highest magnitude open chromatin features

that left primarily promoters, not enhancers, for the method to prioritize (**Supplemental Text**). Notably, a high performance, runner-up method (Aerts CREsted) employed deep learning models, trained with the CREsted package¹⁸, that learned to predict open chromatin from DNA sequence. Sequence models are of particular interest since they have the potential to identify TF motifs that make up cell type-specific enhancer codes^{19,20}, including repressive elements.

Meta-analysis of performance across teams and cell types

To assess whether teams identified similar or distinct enhancers, we computed pairwise intersections of submissions from the top-performing methods (**Fig. 2A**). Methods with similar performance predicted similar enhancers, and the top three methods (Aerts scATAC_{triplet}, ArchR and PeakRankR) had the highest agreement. A majority of On-Target enhancers were identifiable across many approaches (**Fig. 2B**), yet some were not recovered by any method, likely due to low ATAC-seq signal (**Supplemental Fig. 3**).

The Aerts, PeakRankR, and ArchR methods consistently included On-Target enhancers in the top rankings (**Fig. 2C**), while the Aerts method deprioritized more Mixed-Target, Off-Target and No-Labeling (**Fig. 2D**). Interestingly, compared to the top-performing methods, methods that used DNA sequence (Aerts CREsted) and HiC (Gillis) data better prioritized enhancers for L6b neurons, and the CREsted method better prioritized enhancers for low-abundance Sst Chodl inhibitory neurons (**Fig. 2E, Supplemental Figs. 4-5**). In addition, Aerts scATAC_{triplet} and Aerts CREsted were better than ArchR and PeakRankR at deprioritizing lower quality enhancers for most cell types (**Supplemental Figs. 6-7**).

Next, we compared performance in prioritizing L5 ET enhancers, the largest set of validated enhancers. The top five strongest and most specific On-Target L5 ET enhancers were ranked highly by most methods, with somewhat lower rankings from Gillis (**Fig. 2F**). Most methods correctly scored a strong L5 ET enhancer AiE0456m higher than a weak enhancer AiE0460m, likely due to higher ATAC-seq signal in AiE0456m and the presence of multiple *Pou3f1* motifs (**Fig. 2F**), a canonical TF of L5 ET neurons¹. The strong enhancer AiE0463m had low ATAC-seq signal but multiple *Pou3f1* motifs and was correctly prioritized only by the Aerts CREsted model (**Fig. 2F, Supplemental Fig. 7**). Thus, methods that can learn from DNA sequence have an advantage to prioritize strong and specific enhancers that are associated with chromatin regions with limited accessibility.

Additional genomic features can help predict enhancer activity

Since chromatin accessibility sometimes failed to predict cell type-specific activity, we tested if prediction accuracy could be improved by including additional enhancer features: (1) H3K27ac, a histone modification found at active enhancers²¹ from mouse cortical Paired-Tag data^{13,22}, (2) Activity-By-Contact (ABC) scores, the product of chromatin accessibility and contact frequency from HiC^{13,23}, and (3) conservation of chromatin accessibility between human and mouse¹³. Similar to chromatin accessibility, cell type-specific H3K27ac and ABC scores were significantly higher for On-Target validated enhancers compared to Mixed-Target, Off-Target, No-Labeling and a random control (**Fig. 3A**). On-Target enhancers had significantly higher conservation of accessibility (**Fig. 3A**) and sequence (**Supplemental Fig. 8**) compared with No-Labeling and Random but not Mixed-Target or Off-Target enhancers. Thus, conservation of accessibility predicted overall enhancer activity, while H3K27ac and ABC predicted cell type-specific enhancer activity.

For validated enhancers, cell type-specific chromatin accessibility was notably correlated with cell type-specific H3K27ac ($r = 0.57$) and ABC scores ($r = 0.38$), but not epigenetic conservation ($r = 0.07$) (**Fig. 3B**,

Supplemental Fig. 9). For example, On-Target enhancer AiE2121m had high chromatin accessibility and H3K27ac specifically in astrocytes (**Fig. 3B,C**). In contrast, No-Labeling element AiE0358h had high chromatin accessibility but not H3K27ac specifically in astrocytes (**Fig. 3B,C**). These examples demonstrate a potential for H3K27ac signal to improve the accuracy of enhancer predictions by distinguishing functional from non-functional chromatin accessible candidate enhancers. Finally, we trained a binary random forest classification model to predict on-target enhancers from all others. In the held-out test set, this model resulted in an AUC of 0.75. This model is better able to identify poor enhancers (precision 0.77, recall 0.80) than on-target enhancers (precision 0.52, recall 0.47). While the precision is relatively low, this model was better able to identify on-target enhancers than methods used to initially identify them (31% of mouse enhancers are in the on-target class). In this model open chromatin metrics (ATAC-seq strength or z-score and cell type specificity), linear sequence conservation (% matching bps) and GC content were the most informative features (**Fig. 3D, Supplemental Fig. 10**).

Re-scored enhancers validate model predictions

To investigate differences in predicted versus *in vivo* enhancer activity, we manually inspected the ATAC-seq signals and prediction and nucleotide contribution scores from the CREsted models for all 677 tested enhancers. We classified each enhancer into one of five categories: ‘Explainable positives’ (48%) have *in vivo* activity and high ATAC specificity and CREsted scores in the corresponding cell type; ‘Explainable negatives’ (25.4%) have no *in vivo* activity and low ATAC or CREsted specificity; ‘Unexplainable positives’ (6.2%) have activity but low ATAC and CREsted specificity; ‘Unexplainable negatives’ (13.9%) have no reported activity yet high ATAC and CREsted scores, suggesting potential false-negative experimental results; and ‘Missing data’ (6.5%) for the remaining enhancers. We visualized enhancers based on the peak-scaled ATAC signal and labeled by the targeted cell type (**Fig. 4A**). Explainable On-Target enhancers were well segregated and overlapped with some unexplainable No-Labeling (negative) enhancers. This suggested that some negative enhancers may have weak *in vivo* activity that was missed in the initial evaluation (i.e., experimental false negatives).

Therefore, we re-evaluated all 267 No-Labeling enhancers and found that 66 weakly drove SYFP2 expression in the brain (**Fig. 4B**). 52 of 66 enhancers had weak expression in the targeted cell type and were rescored to On-Target. These enhancers were enriched for unexplainable versus explainable negatives, which supports our predictions of enhancer activity. Interestingly, among the 140 explainable negatives that were confirmed to have no activity after rescoring, 45.7% had ATAC-seq signal in their targeted cell type but no support from the CREsted model. Conversely, only two enhancers (1.3%) contained a specific CREsted prediction and no ATAC-seq peak. The remaining 74 enhancers (53%) were not predicted to have cell type-specific activity by either ATAC-seq or CREsted. Thus, considering DNA sequence along with ATAC-seq signal can help avoid testing inactive enhancers.

To illustrate the rescoring process, we show two enhancers that had scATAC-seq, CREsted and H3k27ac scores that were specific to oligodendrocytes (Oligo). First, an On-Target Oligo enhancer was predicted by the CREsted model to include two important candidate TF binding sites: one for the established Oligo TF *Sox10*^{24,25}, and one for the Ap-1 TF group, known to cooperate with cell type-specific TFs for enhancer function²⁶ (**Fig. 4C**). Second, a No-Labeling enhancer was rescored as weakly On-Target for Oligos, and this was consistent with predictions from all modalities and a SOX10 binding site (**Fig. 4D**). Hence, insights into cell type enhancer codes enabled prediction of functional enhancers and their relative levels of *in vivo* activity.

Using the rescored enhancers, we assessed the multi-label classification performance of enhancer activity by calculating precision and recall at different score thresholds. ATAC-seq modalities scored higher (average precision, AP = 0.47 for count-normalized pseudobulk scATAC and AP = 0.52 for peak-scaled scATAC) than the CREsted sequence model (AP = 0.41) and H3k27ac (AP = 0.26) (**Fig. 4E**). A receiver operating characteristic (ROC) curve highlights the same trends for the different modalities (**Supplemental Fig. 11**). Interestingly, combining outputs from the CREsted and scATAC-seq models improves prediction of *in vivo* enhancer activity (AP = 0.54 with peak-scaled ATAC, AP = 0.52 without peak-scaling), demonstrating that some cell type-specific regulatory sequences were learned.

Next, we examined if the CREsted model captured sequence differences that were associated with the magnitude of *in vivo* activity of On-Target enhancers. On average, the model predicted higher scores for strong versus weak enhancers for 9 out of 14 cell types (**Fig. 4F, Supplemental Fig. 12**). Additionally, we investigated how well models identified explainable negative enhancers. The CREsted model significantly ($P < 0.05$) outperformed scATAC approaches in avoiding false positives (**Fig. 4G**) and false negatives (**Supplemental Fig. 13**). Overall, these analyses underscore the benefits of using sequence models to improve non-functional enhancer prediction, prioritize strong enhancers, and lay the groundwork for understanding cell type enhancer codes.

Discussion

The BICCN community enhancer challenge provides a valuable benchmark for future computational methods aimed at predicting functional cell type-specific enhancers. With 677 enhancers validated on a standardized pipeline and scoring criteria, this represents the largest collection of its kind⁶. Importantly, both data and code are publicly available, facilitating further research and method refinement. However, limitations exist, including use of bulk ATAC-seq datasets and selection of enhancers near cell type markers that may not represent the most specific peaks per cell type. In addition, the epifluorescence scoring, while valuable, is an imperfect estimate of target specificity based on cell morphology clues and spatial distributions of labeled cells. Finally, SSv4 data was collected from mouse V1 and may not always align with model predictions based on mouse M1 multi-omic data, although cell subclasses share similar molecular profiles across the mouse neocortex^{7,27}.

The validation data from this challenge reveal which features are critical for predicting enhancer function, and the highest-performing methods primarily leveraged the specificity of ATAC-seq peaks in the mouse data set. Interestingly, the top submission combined RNA-seq with ATAC-seq to predict cell type-specific TF-enhancer-gene triplets. Additionally, incorporating HiC data was valuable for predicting enhancers in specific neuron types, such as L6b. H3K27ac and ABC scores tended to be higher in On-Target enhancers, although with lower recall and precision compared to cell-type specific ATAC-seq signals. Moreover, biological priors played a crucial role in the success of these models. For instance, including moderately sized ATAC-seq peaks was critical to predict candidate enhancers since the largest peaks often represent promoters.

Despite initial expectations that deep learning and DNA sequence models would outperform, the results showed that simpler ranking of ATAC-seq peaks by their cell type specific signal performed similarly. However, the top model did not consistently excel across all cell types and enhancer categories. The distinct methodologies used by the top three teams indicate that there are opportunities to refine enhancer prioritization. For example, the CREsted sequence model avoided Mixed-Target or Off-Target enhancers and predicted a strong oligodendrocyte enhancer by finding a coactivating TF motif AP-1 near a motif for *Sox10*, an oligodendrocyte marker^{24,25}. Additionally, sequence models outperformed ATAC-

seq only models in identifying Sst Chodl peaks, where the latter model may have struggled due to few cells being profiled from this rare type and thus a low ATAC-seq signal-to-noise ratio.

Looking forward, a major goal of the field is to develop a comprehensive toolkit to target cell types across the brain. Improving data coverage and quality is crucial, as many On-Target enhancers were missed due to low ATAC-seq signal. Fortunately, on-going work supported by the NIH BRAIN Initiative is focused on whole-brain atlasing of cell types, including high read-depth single cell ATAC-seq, HiC, and histone marker profiling. Sequence models will enable the rational design of enhancers tailored to cell types or groups of types, a strategy successfully applied in fruit fly models²⁰. Cross-species sequence models were under-represented in this challenge, but On-Target enhancers tend to have conserved open chromatin and sequence, and TF regulatory networks show conservation across primates and rodents¹. Additionally, sequence models trained on ATAC-seq data from multiple species enhance the prediction of chromatin accessibility¹³. Given these observations, we propose that decoding conserved enhancer codes presents a promising avenue for developing robust and precise tools to target cell types across many species.

Expanding validation experiments to include a broader range of brain regions and cell types, especially those with few specific markers will be essential. Community sharing of raw data will enable re-evaluation with future algorithms and comprehensive whole-brain analysis, while negative results will provide valuable insights into DNA repressor codes and enhancer function. Cross-species testing, including in non-human primates, will help assess the predictive power of these models across model organisms and will bolster confidence in translational applications for humans.

Integrating modeling and experimental approaches will be key to advancing enhancer tool development. Selecting enhancers that offer the most informative data for models will refine predictions and improve success rates. Testing enhancers with conflicting predictions from different modalities, such as ATAC-seq versus DNA sequence, could yield critical insights, even if they do not result in the most effective tools. Additionally, reinterpreting experiments with no labeling that have strong model support may uncover challenges in transducing some cell types such as has been reported for microglia²⁸. Ultimately, we need interpretable models that help to generate viral tools to target cell types and provide insights into cell type identity and genetic regulation in the context of human evolution and disease.

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Author contributions:

Viral tool testing: BPL, BT, DD, JKM, JTT, MH, TLD, YB

Data analysis: BL, DA, DD, DM, ECE, EJA, FW, GH, GP, JDM, JG, JTT, KZ, MH, NH, NJJ, NK, NRZ, RL, SA, SDW, SS, TEB, VK, YB, YT

Data interpretation: BL, BR, BT, DA, DD, DM, ECE, EJA, ESL, FW, GH, GP, JDM, JG, JM, JRE, JTT, KZ, MH, NH, NJJ, NK, NRZ, RL, SA, SDW, SS, TEB, VK, YB, YT

Writing manuscript: BL, BT, DD, FW, JG, JTT, KZ, MH, NJJ, NK, NRZ, RL, SA, SS, TEB, YB, YT

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Data and materials availability: Cortical cell type AAV-based tools are described in Ben-Simon et al.⁶ and are available from Addgene (<https://www.addgene.org/collections/brain-armamentarium>).

Methods

Single nucleus molecular profiling

10x multiome ATAC + Gene Expression and methyl-3C-sequencing (snm3C-seq) experiments were carried out on the same tissue samples from human, macaque, marmoset, and mouse M1, as described¹³.

In vivo enhancer validation data

Detailed experimental methods are described in the companion paper⁶ and other recent viral tools publications^{4,5,25}.

Primary screen scoring

Each enhancer vector was screened and scored based on the labeling pattern it produced across the entire brain, with additional emphasis on cortical populations. First, each region of the brain where labeling of cell somata was observed was manually scored based on the labeling brightness and density, classifying each into either low or high. In addition, we created 11 categories of cell populations within the neocortex that could be visually distinguished one from the other. Whenever labeling was observed in one or more of these cortical populations, each population was individually evaluated based on its own brightness and density. Whereas brightness was classified based on whether the labeling was stronger or weaker than the common brightness observed across all experiments, density was evaluated based on the expected density of cells for each of the scored regions or populations, using the nuclear markers as reference. To determine target specificity, we aligned each target cell population with the labeled population which best matches its known anatomical location, distribution, and morphological characteristics. We determined an enhancer to be “On-Target” if the target population aligned with the labeled population, “Mixed-Target” if labeling was observed in populations other populations, in addition to the target one,

“Off-Target” if labeling was observed exclusively in population/s other than the target population, and “No-Labeling” if no labeling was observed in the neocortex, regardless of whether labeling was observed in other brain regions.

Cell type enhancer activity quantified by single cell RNA-seq

SSv4 data for cortical enhancers were generated from mouse primary visual cortex (V1/VISp) and mapped to the Allen Institute AIT2.1.1 VISp taxonomy as described in Ben-Simon et al. ⁶. The data were reused in this study to establish ground truth cell type specificity in combination with primary screening expression analysis.

Benchmark metrics for evaluation of enhancer prioritization methods

We defined an interpretable benchmark metric based on the experimentally validated enhancer collection reported in Ben-Simon et al. ⁶. This benchmark metric is available to the community at: <https://github.com/AllenInstitute/EnhancerBenchmark>. The benchmark metric is a composite of an epifluorescence imaging score and Ssv4 quantifications score per enhancer that captures different properties of enhancer activity. Epifluorescence validation of an enhancer captures broad patterns such as cortical layer labeling and provides cell type-specificity when cellular morphologies are known. Additionally, the intensity of SYFP2 fluorescence provides a measure of enhancer strength. We scored each validated enhancer based on the cell type-specific rank and placed more weight on strong enhancers being enriched in the top ranks. However, the epifluorescence validation does not provide a quantitative measure of specificity for each cell type targeted by the enhancer virus. To achieve this, we used Ssv4 to quantify the transcriptome of cells with high SYFP2 fluorescence and mapped these cells to a cortical taxonomy to determine accurate abundances of each cell type targeted by the enhancer.

The epifluorescence metric was designed to be minimized when teams arranged On-Target enhancers in the top ranks per-cell type and Mixed Target enhancers relatively lower down the ranks as these enhancers both target the intended cell type as well as additional unintended cell types. Negative enhancer categories including Off-Target and No-Labeling enhancers optimize the metric when placed at the bottom of the ranked lists per-celltype. For each validated enhancer $E_i \in E_{validated}$ in the ranked list across cell types we multiply the predicted rank R_{E_i} of the enhancer with an indicator variable $I_{Category}$ which encodes enhancer categories (On-Target, Mixed-Target, Off-Target or No-Labeling). Then the metric is weighted by an indicator variable $I_{Strength}$ encoding the strength of SYFP2 epifluorescence (Strong, Weak or None).

$$Epi\text{fluorescence}_{metric} = \sum_{E_i \in E_{validated}} (Rank_{E_i} * I_{Category_{E_i}}) * I_{Strength_{E_i}} \quad (1)$$

where:

$$I_{Category} = \begin{cases} -1 & \text{On-Target} \\ 0.5 & \text{Mixed-Target} \\ 1 & \text{Off-Target|No-Labeling} \end{cases} \quad I_{Strength} = \begin{cases} 2 & \text{Strong} \\ 1.5 & \text{Weak} \\ 1 & \text{None} \end{cases}$$

The Ssv4 metric was designed to be minimized and replaces epifluorescence strength $I_{Strength}$ with a quantification of cell type-specificity computed the fraction of cells labeled as the intended target cell type $SSv4_{E_i, celltype_{target}} [0 - 1]$. We then compute the Ssv4 metric for each validated enhancer $E_i \in$

$E_{validated}$ by multiplying the predicted rank R_{E_i} with the associated category $I_{Category}$ and the fraction of cells labeled as the targeted cell type $SSv4_{E_i, celltype_{target}}$.

$$SSv4_{metric} = \sum_{E_i}^{E_{validated}} (Rank_{E_i} * I_{Category_{E_i}}) * SSv4_{E_i, celltype_{target}} \quad (2)$$

The composite benchmark score is then computed as the unweighted summation of Epi_metric and SSv4_metric.

$$Benchmark_{metric} = Epi_{fluorescence_{metric}} + SSv4_{metric} \quad (3)$$

To define a normalized benchmark score ranging between 0 (worse) and 1 (better) we created an optimally ranked enhancer list per cell type and computed the associated benchmark metric. Then the benchmark metric score is divided by this optimal score and subtracted from 1 to achieve a normalized metric with the desired directionality.

To prevent overfitting during the challenge, we held back enhancers targeting excitatory cell types to ensure that no team could gain an advantage by learning patterns in the hidden enhancer validation data over the course of multiple challenge submissions.

Recovery curves of functional enhancers per method

To assess the recovery of validated cell type specific enhancers based on the cell type specific rankings of each submission a recovery approach was used. In brief, the set of ground truth cell type specific enhancers was defined as the genomic regions (candidate enhancers) for which the specificity was classified as "On-Target". Mouse genomic coordinates were used for candidate enhancers originating from the mouse genome, and coordinates lifted over to the mouse genome were used for candidate enhancers originating from the human genome. The genomic regions in each cell type specific ranking were intersected with the ground truth cell type specific enhancers using pyranges identifying hits along the ranking (genomic regions in the ranking overlapping with multiple ground truth enhancers were only counted once). Recovery curves were drawn for each submission by calculating the cumulative sum of the union of hits along all cell type specific rankings per cell type. In cases where the ranking was shorter than 10,000 elements, the ranking was padded with non-hits up to a length of 10,000 elements. Normalized enrichment scores (NES) per cell type specific ranking per submission were calculated as the area under the curve of the recovery curve (AUC) up to the 1,000th element and dividing this by the AUC up to the 1,000th element of the average recovery curve of 100 random rankings.

Manual annotation of validated enhancers

We manually inspected a subset of the validated regions by classifying them into five categories: explainable positives, unexplainable positives, explainable negatives, unexplainable negatives, and undetermined. Explainable positives are defined as functional (both 'strong' and 'weak', and both 'On-Target' and 'Mixed-Target') enhancers that have either a strong accessibility peak in the scATAC-seq data, a strong prediction from the CREsted model, or both, in their intended target cell type(s). Unexplainable positives are functional enhancers that do not have a clear peak and/or prediction in their target cell types, but still show activity for those cell types, or they do have a strong peak and/or prediction in their targets, but show activity in other cell types. For validated enhancer candidates that did not show

functionality (negatives), we identify explainable negatives as regions that either do not have a specific peak, a specific prediction, or both, in the target cell type. Unexplainable negatives have either a peak, prediction, or both in their target cell types, as well as the presence of positive contribution scores in one or more motifs obtained from the CREsted model. Undetermined regions do not contain enough decisive information to classify them into one of the other four categories, often because of their target cell types not being included in the original dataset.

Precision-recall curves of different modalities

To compare the different modalities directly on the 677 validated enhancers, we scored each enhancer per modality by taking per cell type the score over the sum of all scores, scaled by the absolute value of the score. We generated a target binary matrix based on the on- and Mixed-Target labels, combined with the SSv4 results, to label the targeted cell types on enhancer functionality, and calculated per cell type the precision and recall over different prediction thresholds. For the scATAC-seq and H3k27ac data, we took the average counts over the exact region, for the CREsted model the enhancer regions were put in the center of a 2114 bp background region, since the model takes in a fixed 2114 bp sequence.

Random forest modeling using biological priors and per enhancer genomics measures

To examine the importance of additional variables determining enhancer specificity, we modeled per enhancer metrics, including distance to the nearest transcription start site, mouse-human enhancer sequence conservation, and published ChIP seq data. ChIP seq data that overlapped enhancers were found using the GenomicRanges package in R, and all overlapping bins for each enhancer were summed to get values used for downstream analysis. Random forest models were constructed using scikit-learn version 1.3.0. The “% matching bps” measure was determined by calculating the percentage of the initial mouse enhancer length that is preserved upon examination of the best BLAST alignment with the human genome. For random forest model development, we used a 70/30 split for training and a held-out test set, respectively. Prior to testing on the held-out test set, models were constructed and validated using 10 fold cross-validation using GridSearchCV. The importance and statistical significance of variables is reported in Figures 3D and S10. Statistical significance was calculated using ANOVA, followed by a Tukey post hoc test.

Stein Aerts lab methods:

ATAC-based rankings

We investigated the performance of rankings purely based on scATAC-seq data. For all consensus peaks, we obtained the mean accessibility per cell type from the pseudo bulked cell type-specific accessibility tracks. We ranked regions based on the Gini index of their accessibility profile over all cell types to obtain the highest and most specific peaks per cell type. Using the Gini index only provides one value per region, so we assigned that value to the cell type with the highest peak value and gave a zero score to all the other cell types in that region. To further optimize this approach, we augmented the scATAC-seq data by merging the provided mouse dataset with publicly available mouse motor cortex datasets^{17,29}. We merged datasets by manually matching corresponding cell types, and weighted peak heights across datasets by using the number of cells per matched cell type.

Normalization of peak heights across cell types

To create peak-scaled scATAC tracks across cell types, we used the CREsted peak normalization functionality with default parameters. In short, the top 1% (based on peak strength) of peaks per cell type

are taken, and based on those, scalars are created which are multiplied with the peak heights of their corresponding cell types to put peaks across different cell types in a more comparable range.

Sequence-based deep learning models

We trained two types of convolutional-based sequence-based accessibility prediction models from the CREsted package. The first is a peak regression model, a 9-layer convolutional neural network (CNN) model, trained on scATAC-seq data for chromatin accessibility analysis. It uses 2,114 bp DNA sequences as input and predicts the average accessibility signal per cell type on the center 1,000 bp of that region, inspired by ChromBPNNet³⁰. A second approach is topic classification, which was the modeling method used in previously published models^{9,31–33}. We analyzed the scATAC-seq data using pycisTopic¹⁷ to obtain topics per region required for training such models. Transfer learning to differentially accessible regions (DARs) made it possible to obtain cell type-specific predictions, as was done in Hecker et al. 2024⁹.

We used two methods of ranking all the consensus peaks for these models. The first one was using the Gini index on the prediction scores over the cell types per region. The second one was calculating per region, for each prediction per cell type, the product of the difference between a given cell type prediction and the highest prediction in any other cell type, and the ratio between them. This gave a specificity ranking per cell type, per region, which could be sorted to generate global rankings.

Pattern based enhancer scoring

We reasoned that regions with heterogeneous motif content were more likely to be functional enhancers. For this purpose, we generated SHapley Additive exPlanations (SHAP) values for the top 10,000 regions per cell type from DeepBICCN, generating explanations for the prediction score of that cell type. Of these, the top 5,000 regions per cell type were used to identify recurring, important patterns using the tfmodisco-lite package³⁴. To rank regions based on motif diversity, we calculated the Shannon diversity index based on the pattern hits. This index was multiplied by a signal over noise metric that was defined, per region, as the number of seqlets (short stretches of DNA with a high importance score) having a pattern hit divided by all seqlets.

Scenic+ triplet scores

We ran SCENIC+ on the provided mouse multiome dataset (scRNA-seq + scATAC-seq) using default parameters¹⁷. Based on these results we generated a ranking of all transcription factor (TF)-region-gene triplets, which is the aggregated ranking (see below) of the TF- and region-to-gene importance scores and TF-to-region ranking based on the motif score of all motifs annotated to that TF. To make this ranking cell type-specific we combined it with the gini-based cell type-specific ATAC ranking.

Aggregate rankings of multiple methods

To combine multiple rankings of different methods, we used OrderStatistics³⁵. In brief, rankings were generated based on the score of each method, ties were broken by assigning incremental rankings to tied regions based on the order in which they happen to occur. Next, rank-ratios were calculated as the ranking divided by the number of regions in each ranking and combined in a single ranking using the formula described in [ref] and implemented in the SCENIC+ package¹⁷.

Jesse Gillis lab method:

We employed a multi-modal approach to predict cell-type-specific enhancers in the mouse brain, utilizing three distinct data types: single-cell RNA sequencing (scRNA-seq), single-cell Assay for Transposase-

Accessible Chromatin sequencing (scATAC-seq), and meta-Hi-C. For scATAC-seq data, we obtained binarized MACS2 peak scores from three publicly available sources^{7,13,29}. These scores were then z-scored at each genomic bin to capture cell-type specificity accurately. To enhance the power of scATAC-seq data further, we aggregated the binarized peaks from the three scATAC-seq sources to create a robust scATAC-seq matrix. Again, this matrix was z-scored at each bin to enhance cell-type specificity.

In the case of meta-Hi-C, we used our in-house method to obtain cell-type-specific contact vectors by combining meta-Hi-C data with scRNA-seq marker genes. Previously, we aggregated hundreds of Hi-C contact matrices for mouse Hi-C to generate meta-Hi-C maps available at <https://labshare.cshl.edu/shares/gillislab/resource/HiC/>³⁶. Similarly, we utilized scRNA-seq data to obtain reliable markers for each brain cell type in mice³⁷. The cell-type-specific profile was derived by computing the mean of the top markers for each cell type and smoothing the Hi-C contact vector using various marker subsets. The resulting meta-Hi-C matrix was z-scored at each bin to enhance cell-type specificity. Only intra-chromosomal contact matrices were used for this analysis. The Hi-C vectors are slightly updated from the time of submission to the challenge, however this does not alter the validation results significantly.

To integrate the meta-ATAC-seq and meta-Hi-C data, we multiplied the ATAC-seq peak scores with the Hi-C contact scores for each cell type, resulting in the metaATAC X metaHi-C matrix. This matrix was also z-scored at each bin to capture cell-type-specific enhancers accurately.

Saroja Somasundaram lab method:

PeakRankR is an R package (<https://github.com/AllenInstitute/PeakRankR>) for prioritizing and ranking cell type specific peaks given chromatin accessibility data. It uses a systematic approach to sum the effects of positive features for a given peak in a cell type and rank them. The features computed by PeakRankR include: (1) Specificity is computed using an R version of `multiBigWigSummary` from `deeptools` (<https://deeptools.readthedocs.io/en/develop/>); (2) Sensitivity is computed by determining the number of cell types which have a peak at the enhancers genomic coordinates; (3) Magnitude, is computed using MACS2 to determine the abundance of the ATAC-Seq signal at the enhancers genomic coordinates for the cell type of interest. Using these three features PeakRankR produces a score per peak as a weighted linear sum of the three features:

$$PeakRankR_{score} = w_{specificity} Specificity_{Peak_i} + w_{sensitivity} Sensitivity_{Peak_i} + w_{magnitude} Magnitude_{Peak_i}$$

where w stands for the weight of each feature. By default each weight variable is set to 1 indicating equal importance for all three features..

We extended PeakRankR to include features that describe the shape of a peak which include: (1) Skew, which measures the asymmetry of read pileups in an enhancer with symmetry (skewness close to 0) ranked higher; (2) Kurtosis, which measures the reads dispersion between center and tails in an enhancer with higher ranked better; (3) Modality, which describes the number of peaks in an enhancer with unimodal ranked higher. These features were calculated using the functions from `modes` (<https://www.rdocumentation.org/packages/modes/versions/0.7.0>) package in R.

Many peak ranking methods require substantial computational resources, and the selected peaks are not always robust. Hence, there is a need for a simple and efficient function to rank peaks that can be cloned to target cell types. PeakRankR helps bridge this gap by extracting the optimum features given a peak set for a cell type and calculating a score depending on specificity and sensitivity.

Yoshiaki Tanaka lab method:

cisMultiDeep is the repository of R and Python scripts and command lines that identify functional CREs from single-cell multi-omics profiles. Given high conservation of the cell type-specific genes, we first obtained an orthologous gene list from Biomart (<https://ensembl.org/info/data/biomart/>)³⁸. On the other hand, the peak conservation was defined by UCSC LiftOver function in KentUtils and Bedtools (v2.30.0)³⁹. Subsequently, the cell type specificity in each gene and peak was estimated in RNA, mCG, mCH, and ATAC profiles by Wilcoxon rank-sum test. Here, the cell type specificity was calculated only in orthologous genes, whereas both conserved and non-conserved peaks were used for the assessment of the cell type specificity. In each cell type, the top 1,000 genes and 10,000 peaks were selected for subsequent deep learning analyses.

To ask if the selected genes/peaks are sufficient to define the cell types, we employed automatically-tuned deep neural network that was designed by Tensorflow python library (v2.9.0) with Keras Tuner API (v1.1.2)^{40,41}. Briefly, at first, dimensionality of the input data (RNA, mCG, mCH, and ATAC profiles) was reduced into 200 by principal component analysis (PCA). Then, the sequential neural network model was built with seven tunable hyperparameters: i) the number of layers (2 to 10 with increment of 1), ii) the number of nodes in hidden layers (50 to 500 with increment of 50), iii) dropout rates (0 to 0.5 with increment of 0.1), iv) activation functions (e.g. sigmoid), v) optimizers (e.g. Adam), vi) learning rate (e.g. 1e-1, 1e-2, 1e-3, 1e-4, 1e-5), and vii) loss functions (e.g. mean squared error loss function). Once the neural network model was optimized, mean absolute SHAP value, which represents the impact of each gene or peak on the cell type determination, was calculated by DeepSHAP that is a technique that can handle the complex and non-linear interactions across features and is optimized to calculate SHAP value for deep neural network⁴².

Chromatin looping enables distal CREs to contact their target genes. Here, we hypothesized that the cell type-specific CREs are physically contacted with various cell type-specific genes. Thus, we ranked the peaks by the sum of the mean absolute SHAP values of the contacted genes by Hi-C loop. If the peak is conserved, we also added up the sum of the mean absolute SHAP values in other species.

Kai Zhang lab method:

Using the genome annotation downloaded from Gencode, for each gene we extracted the 196,608 bp DNA sequences centered around its TSS. We then applied the Enformer model¹⁹ to these sequences to derive sequence embeddings, followed by attention pooling to further reduce the dimensionality. Consequently, the processed sequences were represented as 896 vectors, each with 48 dimensions, corresponding to 128-bp segments of the initial sequence. Additionally, we quantified the ATAC-seq fragments overlapping these segments, adjusting for reads per kilobase million (RPKM).

To identify candidate enhancers, we developed a machine learning framework that integrates ATAC-seq signals and the aforementioned sequence embeddings to predict gene expression profiles. We first transformed normalized counts of ATAC-seq fragments into four-dimensional vectors using convolutional

and self-attention layers. These ATAC embeddings were then merged with sequence embeddings and fed into a sequence of linear layers aimed at predicting levels of gene expression.

To train the model, we partitioned the data into training, validation and testing sets using an 80:10:10 split. We employed the Adam optimizer with a learning rate of 0.0001 and trained the model over 10 epochs, after which the model's performance plateaued.

To determine the enhancer activity score for a specific region, we modified its ATAC signal by setting its count value to zero. The trained model was then used to make two predictions: one for the original (unmasked) input and another for the modified (masked) input. The enhancer activity score was calculated as the difference in the model's predicted gene expression between these two inputs, serving as an indicator of the ATAC signal's impact on the expression of adjacent genes. For each cell type, the top 10,000 highest scored peaks were chosen for submission.

Figures:

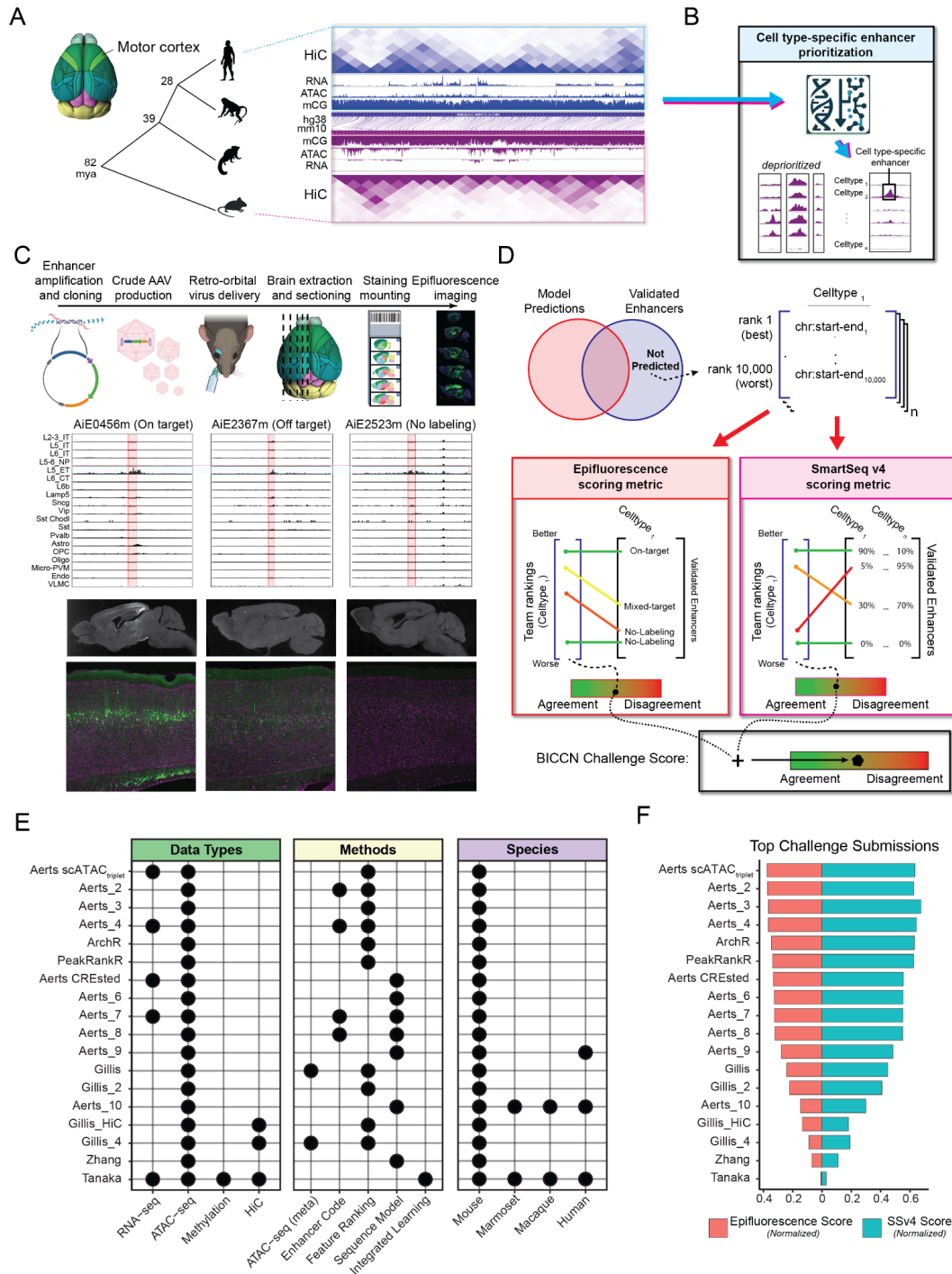


Figure 1. Overview of the enhancer prioritization challenge.

(A) Single nucleus multi-omic data from primary motor cortex of human, macaque, marmoset, and mouse. mya, million years ago.

(B) Schematic of the computational challenge to prioritize candidate cell type-specific enhancers.

- (C) Overview of AAV construction, cell type ATAC-seq specificity, and screening of *in vivo* activity in the mouse brain for three candidate L5 ET enhancers.
- (D) Teams predicted and ranked 10,000 candidate enhancers for each of 19 cortical cell types and were scored based on prioritization of strong, On-Target enhancers.
- (E) Combinations of data and methods for top team submissions.
- (F) Normalized benchmark metrics (**Methods**) based on epifluorescence and SSv4 from *in vivo* screening.

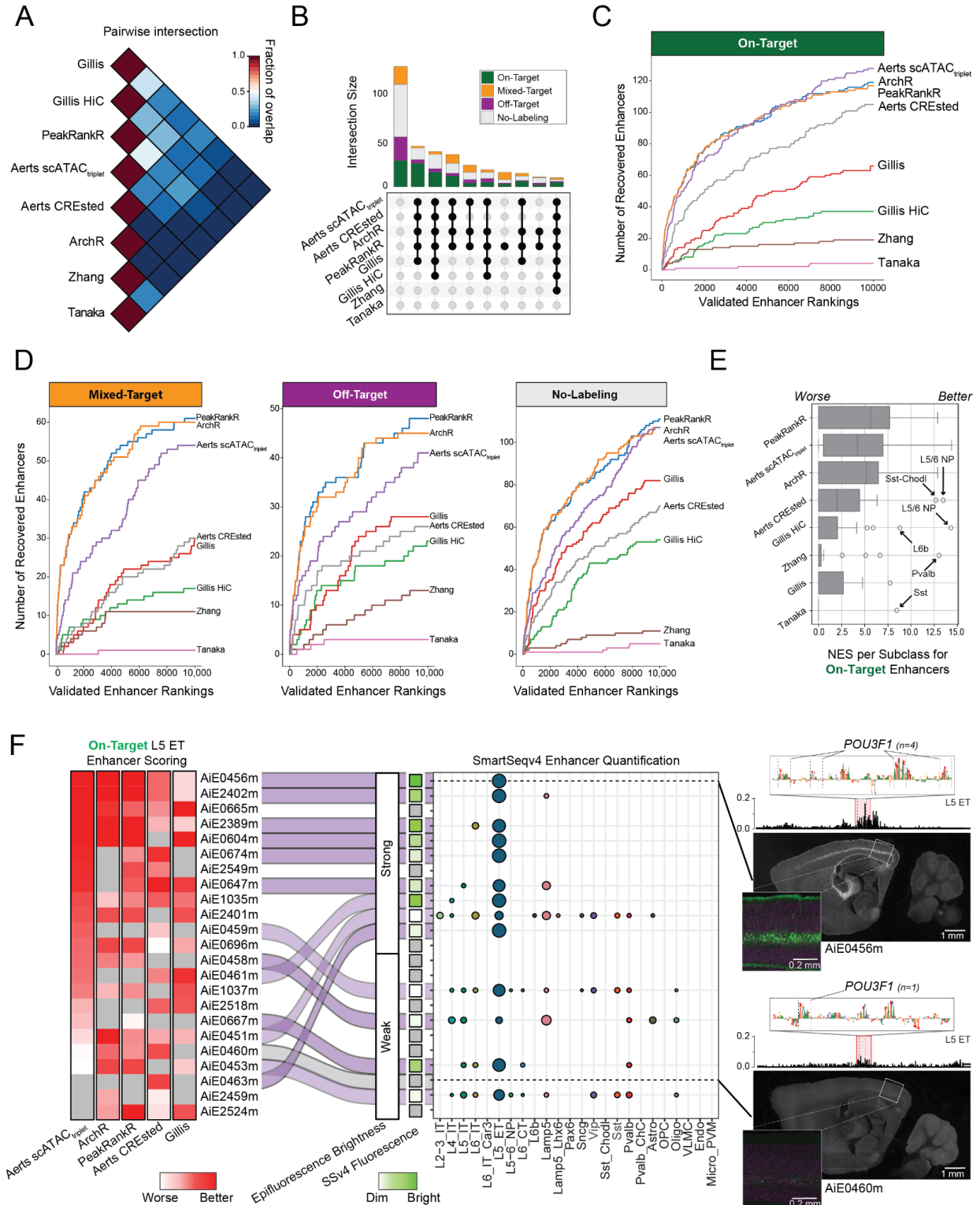


Figure 2. Comparison of team enhancer rankings.

(A) Average proportion of ranked enhancers that overlap between pairs of team submissions for all cell types.

(B) Upset plot showing the number of validated enhancers that were identified by sets of submissions.

(C,D) Rates of identification of (C) On-Target and (D) Mixed-Target, Off-Target and No-Labeling enhancers.
(E) Distributions of normalized enrichment scores (NES) for cell type specific rankings.
(F) Heatmap ordered by Aerts scATAC_{triplet} scoring of L5 ET enhancers and summary of validation results.
Examples of a strong (AiE0456m) and weak (AiE0460m) enhancer with *Pou3f1* motifs identified by the CREsted model in the highlighted region. AiE0456m was also validated with SSv4.

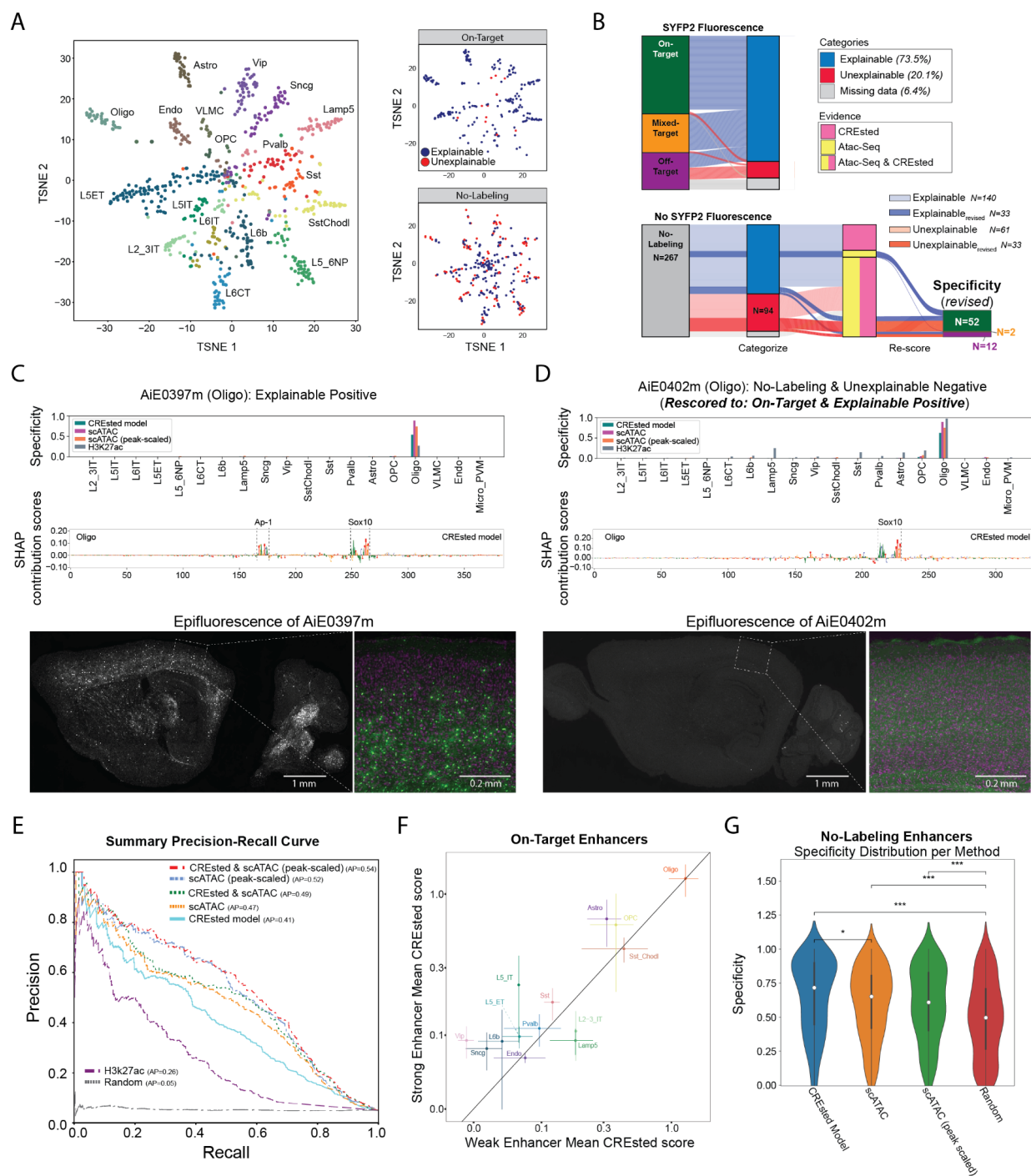


Figure 4. Refinement of models and enhancer screening results.

(A) tSNE plots of enhancers based ATAC-seq specificity and labeled by the targeted cell type. On-Target and No-Labeling enhancers had explainable or unexplainable cell type labeling patterns based on ATAC-seq and DNA sequence (CREsted) model predictions.

(B) River plots of enhancer activity, predictions and rescored of experimental validation data.

(C-D) Model scores, predicted TF motifs and SYFP fluorescence for two Oligo enhancers with (C) strong On-Target and (D) No-Labeling rescored to weak On-Target activity.

(E) Performance of enhancer ranking methods using the rescored enhancer activities. AP, average precision.

(F) CRESTed model scores for strong and weak On-Target enhancers grouped by cell type. Mean +/- SEM.

(G) Comparison of models at identifying No-Labeling enhancers. * $P < 0.05$, *** $P < 0.001$, Wilcoxon rank-sum test, Bonferroni-corrected P-values.

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