SPLASH: A statistical, reference-free genomic algorithm unifies biological discovery

**Graphical abstract**

**Highlights**

- Paradigm for statistical detection of sample-dependent variation in sequencing data
- Computationally efficient, widely applicable, without need for reference genomes
- Finds viral strain mutations, cell-type-specific isoforms, and Ig and TCR diversity
- Finds octopus tissue-specific isoforms, eelgrass/diatom seasonal and geographic variation

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**In brief**

Genomics workflows typically map reads onto a reference genome as the foundation for downstream analyses. However, this poses severe limitations for biological discovery when references are incomplete or nonexistent, and even for intensely studied genomes with rich population-level diversity. SPLASH is a highly efficient framework for statistics-driven analysis of sequence variation directly from raw sequencing data, overcoming previous limitations.

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**SUMMARY**

Today’s genomics workflows typically require alignment to a reference sequence, which limits discovery. We introduce a unifying paradigm, SPLASH (Statistically Primary aLignment Agnostic Sequence Homing), which directly analyzes raw sequencing data, using a statistical test to detect a signature of regulation: sample-specific sequence variation. SPLASH detects many types of variation and can be efficiently run at scale. We show that SPLASH identifies complex mutation patterns in SARS-CoV-2, discovers regulated RNA isoforms at the single-cell level, detects the vast sequence diversity of adaptive immune receptors, and uncovers biology in non-model organisms undocumented in their reference genomes: geographic and seasonal variation and diatom association in eelgrass, an oceanic plant impacted by climate change, and tissue-specific transcripts in octopus. SPLASH is a unifying approach to genomic analysis that enables expansive discovery without metadata or references.

**INTRODUCTION**

Genomics is now foundational to biology, ecology, and medicine, and as sequencing databases grow, so too does the opportunity to leverage them for discovery. How can these data best be analyzed to reveal regulation and function? Traditionally, bioinformatic pipelines start by assigning genomic positions to reads via alignment to a reference genome, an approach with many limitations. For less-studied organisms, references can be partially misassembled, incomplete, or nonexistent. Even in the intensely studied human genome, it was found that understudied populations have large amounts of sequence missing from the current reference\(^1\); such blind spots may exacerbate health disparities. Reference-based methods are not well suited to deal with paralogs and repetitive elements (which comprise \(\sim54\%\) of the human genome\(^2\)), so many analyses simply ignore them. They are also poorly suited for diseases such as cancer that are almost defined by their deviations from the reference and vary even within a single tumor. Additionally, the enormous diversity of viral and microbial genomes and their constant adaptation\(^3,4\) makes it infeasible to define a complete set of references. Practically, alignment to references is computationally intensive, limiting the scale of genomic inference.

When dealing with genomic data, precise statistical analysis is critical. However, alignment-based methods are complex and difficult to model statistically; even seemingly simple tasks, such as calling allele-specific expression, can be fraught with statistical imprecision introduced during alignment.\(^5\) Permutation-based methods are not a panacea; in addition to being slow, they can give 10-fold underestimates of the false discovery rate.\(^6,7\)

Addressing these concerns has led us to a simple, unifying paradigm for statistically detecting signals of biological interest directly from raw sequencing data without using a reference genome, which we call SPLASH (Statistically Primary aLignment Agnostic Sequence Homing). It relies on a simple formalization of sequence variation (short stretches of varying sequences, “targets,” adjacent to short stretches of constant sequences, “anchors”). SPLASH applies to myriad biological questions that can be framed as asking how sequence distributions vary within a set of samples. In the results below, we provide a snapshot of SPLASH’s wide possibilities for discovery, encompassing viral strain variation, single-cell-level alternative isoforms, and antigen receptor diversity in human samples. We also show that SPLASH is easily applied to less-studied organisms: lemur, octopus, and eelgrass. This demonstrates SPLASH’s
potential to discover meaningful sequence variation without the aid of reference genomes across many biological questions and organisms.

RESULTS

SPLASH is a k-mer based, statistics-first approach to identify sample-dependent sequence variation

The goal of SPLASH is to detect sequence variation between a set of samples. SPLASH uses a specific conception of variation based on k-mers, or subsequences of length k, in sequencing reads. This framework leads to a simple but powerful statistical test that identifies variation that is differentially distributed among the samples.

SPLASH only requires that each sample is represented by a separate file of sequencing data (FASTQ). The definition of a “sample” is dictated by the biological question: samples can denote different cells, different tissues, different individuals, or different mixed populations (metagenomics). Samples might differ by conditions as well—different times or treatments and also in other features (cell type, geographic location, etc.); we refer to these generically as “metadata.”

SPLASH characterizes variation using k-mer pairs called anchors and targets (Figure 1A) (k = 27 by default but is adjustable). Every k-mer in the data is an anchor; each k-mer a fixed offset downstream (R, which may be zero) from a given anchor is one of its targets. Note that targets are always defined relative to an anchor. Anchors with more than one target can report on most sequence variations of interest: from changes at a single position to alternative splicing and isoform usage, gene rearrangements, and more.

The SPLASH process is depicted in Figure 1B (detailed in STAR Methods; Figure S1). Conceptually, SPLASH steps through all positions in all reads of all samples, counting all anchor-target pairs. (To decrease computation time, SPLASH can analyze anchors at a subset of read positions, e.g., every fifth position, as used in this work.) SPLASH compiles a counts table for each anchor, with a column for each sample and row for each target; each entry is the count of a given target in a given sample (a contingency table). This requires only one pass through the raw sequences and does not involve reference alignment, so it is computationally efficient. Importantly, we have developed a highly flexible test statistic that captures the desire to find relatively discrete groups of samples with differing variation and controls false positives even for low numbers of observations; it admits a closed-form p value bound, which is thus fast to compute (unlike resampling methods). For each anchor, SPLASH calculates a p value bound for the null hypothesis that the observed target frequencies in all samples come from the same distribution, i.e., that there is no underlying variation of targets between samples. A low p value for an anchor implies that samples differ significantly in which targets they contain.

While SPLASH can use sample metadata (running in a “supervised” mode), SPLASH does not require such information. Indeed, for all the results presented here, SPLASH was run in its default unsupervised mode. For each anchor, SPLASH tries many random splits of the samples and the targets, retains the one that minimizes the p value, and reports the corresponding effect size for this grouping (next paragraph). This process can detect patterned target variation among the samples, if it exists (Figure S3A; STAR Methods).

For each anchor, SPLASH also calculates an “effect size” that ranges from 0 to 1, with 0 meaning that the target distribution is the same between the two groups and 1 meaning that the sets of targets in the two groups are disjoint. Effect size does not require that target distributions of all samples within a group are similar, just that they are different from the other group; thus, effect size can be high even when there are more than two natural groups. Anchors with large effect sizes have target variation that is more discrete across the samples and are more promising for further biological investigation.

To interpret SPLASH’s findings, it can be useful to have longer sequence context than just the anchor and target. Thus, SPLASH also generates a “consensus” for each anchor in each sample. Consensuses are longer sequences assembled from the raw reads of a given sample, looking at every occurrence of an anchor and extending base by base as long as the reads show a consensus (see STAR Methods). Mapping consensuses to protein sequences can identify protein domains—a powerful, reference-free attribution of biological function. Consensuses can also be aligned to sequence databases or reference genomes; aligning only the consensus sequences for significant anchors reduces the typical computational load by over 500-fold compared with the usual approaches that align all reads (Figure S3B).

Figure 1C diagrams some of the differences between traditional alignment-first approaches for SPLASH; Figure 1D outlines some use cases for SPLASH. These guided us in our initial explorations with SPLASH, which are described below. While SPLASH has some adjustable parameters, we did not attempt to tune these (indeed, SPLASH seems robust to a range of parameters; STAR Methods); all analyses were run with the same settings in unsupervised mode (blind to metadata). Despite this, we found that SPLASH performed well across a variety of datasets and, in all cases, found significant patterns of sequence variation (q values for anchors and binomial p values for target-fraction plots are given in the STAR Methods).

SPLASH identifies strain-defining and other mutations in SARS-CoV-2 de novo

Viral genomes have high mutation rates, at the extremes forming quasispecies. The emergence of SARS-CoV-2 was followed by multiple surges caused by variant strains over the course of just 2 years. This is an ideal setting for the application of SPLASH: viruses are always mutating, but out of a sea of mutations, scientists, clinicians, and public health officials want to identify those showing consistent and significant variation.

We applied SPLASH to two SARS-CoV-2 datasets, both viral amplicon Illumina sequencing of nasopharyngeal swabs from infection-positive patients, taken from times when the dominant strains were Delta or Omicron. The samples from South Africa (November to December 2021) represent the rapid rise of Omicron during its first outbreak. The samples from France (December 2021 to February 2022) represent cases of co-infection by more than one strain; the study authors provided as
Figure 1. Overview of SPLASH

(A) An anchor is a sequence of length k (k-mer) in a read; its target is the k-mer that follows it after a fixed offset of length R. An anchor may occur with different targets, which can capture many types of variation; examples are depicted schematically, with the anchor as a blue box and the targets as orange or violet boxes.

(B) SPLASH compiles a table for each anchor, in which the columns are samples, the rows are targets, and the entries are the respective occurrence counts. SPLASH tests multiple random splits of the samples, calculating a test statistic that measures the deviations between each sample’s target distribution and the average target distribution over all samples, searching for the most discriminating split. For the best split identified, SPLASH reports a p value bound.

(C) Alignment-based pipelines are limited by the need for a reference genome, are compute intensive, and are difficult to model statistically because of their complexity. SPLASH detects variation directly from raw reads with rigorous statistics, is compute efficient, and detects many kinds of variation at once.

(D) Some considerations of when SPLASH may be most useful, which reflect its design characteristics.
Figure 2. SPLASH identifies strain-defining and other variations in SARS-CoV-2

In (A)–(C), sets of targets that distinguish SARS-CoV-2 strains are shown; all are in the spike protein (S) gene. Each heatmap has columns for different samples (patients) and rows for different targets; the coloring indicates the fraction of that target observed in that patient. Summary anchor counts are given for rows and columns. Also shown is a map of the categorical metadata of what strains (primary and secondary) were identified in each patient in the original study; these data were not used by SPLASH, but there is evident agreement between the heatmap and the metadata strain assignment. Binomial p value bounds were computed per STAR Methods.

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metadata the assignment of the primary and secondary viral strains for each case (Table S1) (though not used by SPLASH).

SPLASH finds many significant anchors with low q values (<0.05) and high effect sizes (>0.5) directly from sequencing reads (250 for South Africa dataset, 252 for France; Table S2). High effect sizes are expected for anchors whose targets partition samples by strain. To test whether SPLASH recovers strain-defining and other variations, we examined the subset of significant anchors that perfectly map to a reference strain (Original, Delta, or Omicron BA.1 or BA.2; defining mutations taken from CoVariants.org[13]) and call an anchor “strain defining” if it has at least two targets (by definition different) with >5% abundance, at least one of which perfectly matches to a reference strain. We compared the subset with a control set of anchors, those that are most abundant across all the reads. In the South Africa dataset, 98% (126/128) of SPLASH anchors that mapped perfectly were strain defining vs. 7/201 (3.5%) in the control set (hypergeometric p value < 1.7E − 79). In the France dataset, 100% (39/39) of SPLASH anchors were strain defining vs. 8.4% (21/250) of control anchors (hypergeometric p value < 2.6E − 33). Nearly all of the control anchors have only a single abundant target. Thus, SPLASH, though blind to strain reference sequences and sample metadata, detects strain differences with high precision.

Figure 2 shows exemplary strain-defining mutations identified by SPLASH in the Spike protein (S gene). Figure 2A shows an anchor that distinguishes at the major lineage level: one target has no mutations and is consistent with Delta; the other has the mutation K417N, found in all Omicron strains (but not in Delta or Original). Target fractions across cases are consistent with the strain assignment metadata. Figure 2B shows an anchor that discriminates sub-lineages: one target has no mutation, consistent with Delta; the second target has the 3-nt deletion NL211I and the 9-nt insertion R214REPE, which are Omicron BA.1 specific; the third target has the mutation V213G which is Omicron BA.2 specific. Figure 2C shows an anchor that detects emergent mutations not in our references. One target has a pair of mutations, N679K and P681H, found in all Omicron strains. The other targets all have P681R, a Delta-specific mutation, but two targets additionally encode Q677H (by different mutations). Q677 mutations have arisen independently multiple times in different lineages,11,12 and Q677H in several strain backgrounds enhanced infectivity, syncytia formation, and resistance to neutralizing antibodies in pseudotype assay.13

SPLASH results can also be analyzed completely without a reference genome by examining their coding potential. The consensus sequences are translated in silico to amino acid sequences (in all six reading frames) and matched against a database of protein domain models such as Pfam.14 Protein profiles that are more frequently associated with significant SPLASH anchors, compared with control anchors, are candidates for proteins with important patterns of variation. The distribution of protein domains for SPLASH anchors was statistically different from controls (chi-squared test p values: France < 3.7E − 7, South Africa < 2.5E − 39 (Figure 2D). The top four protein domains in both datasets were beta-coronavirus receptor-binding domain (RBD; within S1 region of spike protein), coronavirus S2 domain (within spike protein), coronavirus M protein, and coronavirus ORF7a protein. By contrast, the bottom four domains for each dataset were completely different from each other. Protein profiling pinpoints domains undergoing high variation; for example, in the South Africa dataset, the spike S2 domain had 23 SPLASH vs. 3 control hits, p = 2.9E − 6 (corrected hypergeometric p value). The spike protein is the major site of antigenic variation in coronaviruses, as it is a principal focus of the immune response; the RBD is well known as a target for natural and therapeutic neutralizing antibodies,15 but, in addition, about 50% of natural anti-spike antibodies are directed against the S2 domain.16

We also carried out SPLASH protein domain profiling on an unrelated virus: rotavirus.17 The domains enriched over controls were rotavirus VP3 and NSP3 proteins (Figure S2). These two proteins have roles in blocking host innate immunity.18 Thus, variation in viral protein domains interacting with the immune system may be a recurring theme in SPLASH protein profiling of viral strains.

In summary, SPLASH finds patterns of variation in SARS-CoV-2, including those characterizing strains, without requiring reference sequences or metadata; the methodology should be generally applicable to other viruses. More broadly, SPLASH may be useful in surveillance for new strains or even completely new pathogens and to cluster patients directly from raw sequencing samples.

SPLASH identifies regulated expression of paralogs and HLA in single cell RNA-seq

Current approaches to single-cell transcriptomics are reference based and specialized; we sought to see whether SPLASH’s unifying methodology could also be applied to single-cell sequencing data generated with the Smart-Seq2 (SS2) protocol,19 which provides broad transcript coverage.

Our first testbed was human macrophage and capillary cells from the Human Lung Cell Atlas20 (Tables S2 and S3), as it was recently established that these cell types have regulated alternative splicing in MYL6, a light-chain subunit of myosin motor protein, which serves as a positive control.21 As expected, among

(A) Distinguishing at the major strain level: target 1 (no mutation) matches Delta; target 2 contains K417N, found in all Omicron (both BA.1 and BA.2 sub-strains); two patients co-infected with Delta and Omicron show both targets (p = 6.4E − 07).

(B) Distinguishing at the sub-strain level: target 1 (no mutation) matches Delta; target 2 with V213G is specific for BA.2; target 3 with a deletion (NL211I) and insertion (R214REPE) is specific for BA.1 (p = 1.6E − 13).

(C) Distinguishing non-strain-related mutations: target 1 has P681R, Delta specific; targets 2 and 3 encode Q677H (by different mutations) and P681R; target 4 has N679K and P681H, Omicron specific (p = 4.9E − 12).

(D) Protein domain profiling in SARS-CoV-2. The top four and bottom four Pfam protein domain hits are shown. S1 receptor-binding domain (RBD) and S2 domain show high variation by SPLASH in both datasets. Other Pfam abbreviations: bCoV, beta-coronavirus; CoV, coronavirus; nucleocap, nucleocapsid; N, N-terminal domain; SARS, severe acute respiratory syndrome coronavirus; M, NS7A, NSP1, NSP8, 3b, and NSP10, viral proteins.
SPLASH’s significant anchors are ones reporting on MYL6 alternative splicing (exon skipping or inclusion) (Figure S4). Interestingly, other SPLASH top anchors also involved myosin light chains, MYL12A and MYL12B, two paralogs with highly similar coding regions (95% nucleotide, 98% amino acid identity for human). Nevertheless, SPLASH finds targets that specifically distinguish them, showing that macrophages express more MYL12A, while capillary cells express more MYL12B, reproducible in two individuals (Figure 3A). Little is known about these genes, but they show differential expression in rat tissues, and there is evolutionary conservation in mammals, birds, and reptiles of adjacent MYL12 paralogs within a syntenic region (e.g., human; rat; Gallus gallus, NCBI Gene IDs 396284 and 770011; Chelonia mydas, Gene IDs 102938771 and 102937279). Besides the small number of amino acid differences between the paralogs, there may also be an important functional role for nucleotide sequence differences, as has been demonstrated for another pair of highly similar cytoskeletal paralogs, beta- and gamma-actin.

In the same data, SPLASH finds cell-type-specific expression of genes in the major histocompatibility complex (MHC), known as HLA in humans. HLA is the most polymorphic region in the genome with the most known disease associations; the polymorphism of HLA class I and class II proteins is intimately tied to their function in antigen presentation for adaptive immunity. Because of the high levels of polymorphism, the region is challenging to represent in a reference genome and for alignment pipelines. Five major haplotypes have been identified at the HLA-DRB locus: all contain DRB1, but some contain a second functional paralog, either DRB3, DRB4, or DRB5. DRB1 is highly polymorphic (3,516 alleles, in March 2023); the paralogs somewhat less so, e.g., DRB4 (236 alleles). SPLASH identifies an anchor with targets that distinguish between the highly similar 3’ untranslated regions (UTRs) of HLA-DRB1 and HLA-DRB4 (class II beta-chains) (Figure 3B). Macrophages express mainly DRB1, while capillary cells mainly express DRB4. This pattern is found in two individuals, who carry different alleles at DRB1 and DRB4. Macrophages are “professional” antigen-presenting cells and constitutively express HLA class II. Not all endothelial cells express class II, although most human capillary cells do; endothelial MHC expression may be strongly cytokine dependent. Thus, class II expression in macrophages and capillary cells is likely to be regulated differently.

In one individual (P3), SPLASH finds a remarkable anchor whose two targets report on HLA-DPA1 and HLA-DBP1 (class II alpha and beta chains, respectively), unique among HLA genes in being organized head to head and transcribed in opposite directions. The anchor lies in a sequence shared by DPB1 and a specific isoform of DPA1, while the targets lie in exons exclusive to each; SPLASH consensus sequences confirm opposite directionality as they bridge splice junctions. Macrophages in this individual express exclusively DPB1, while capillary cells express mainly DPA1 (Figure 3C). This pattern may be haplotype specific, as we did not find a similar anchor for another individual (P2).

A final example is SPLASH’s detection of allele-specific expression of HLA-B in T cells (from a different dataset; see next section). The class I gene HLA-B is the most polymorphic of all HLA genes (9,274 alleles), and HLA-B is the gene with the most anchors found by SPLASH in human T cells (Figure 3D). Since these T cells all come from one individual, this indicates substantial variation in expression of this individual’s two HLA-B alleles at the single-cell level (Figure 3D). Different T cells express a wide range of ratios of the two alleles, some cells expressing both alleles, but others expressing almost entirely one allele or the other (well outside the 98% confidence interval of what is expected by the average ratio). This is in keeping with a preprint that found allele-specific expression of HLA class I genes in normal breast epithelial cells.

Overall, SPLASH finds multiple types of variation regulated at the single-cell level, including paralogs, splicing, and alleles. This gives a glimpse into the complexity of HLA haplotype- and cell-type-specific expression patterns, raising the possibility that disease-related HLA alleles might be expressed differently in key cell types compared to other alleles.

**SPLASH identifies B and T cell receptor diversity in human and lemur single-cell RNA-seq**

Adaptive immune receptors for B cells (immunoglobulin or Ig) and T cells (T cell receptor or TCR) are generated combinatorially through V(D)J recombination, and Ig is further diversified through somatic hypermutation. Rearranged sequences are absent from germline reference genomes and cannot be cataloged comprehensively because of their huge potential diversity, empirically estimated at $10^{10}–10^{11}$ for Ig heavy chains. These genomic loci currently require manual curation because of their complexity and repetitive structure, so few species have high-quality annotations. Existing methods to assign V(D)J rearrangements in single cells depend critically on annotations and so may have blind spots. Since SPLASH is designed to identify sequence diversity without a reference, we predicted that it would identify adaptive immune receptors de novo.

We ran SPLASH on 50 naive human B cells from peripheral blood of one individual and separately on 128 CD4+ human T cells of another individual, taken from Tabula Sapiens, a large multi-organ dataset (Tables S2 and S4). As a first reference-free pass, protein profiling found that the domains enriched in SPLASH anchors in B cells were Ig V-set and C1-set (variable-like and constant-like domains); these two domains were also matches in T cells (attributable to TCR) (Figure 4A). Mapping transcript gene names to SPLASH anchors gives a similar picture: Ig light-chain genes (both kappa and lambda) were strongly hit for B cells; HLA-B (discussed above) and TCR genes (both alpha and beta) were most prominent for T cells. These domains are not found among the control anchors (Figure 4A). Significant anchors were also found in Ig heavy chains, though fewer than in light chains. Ig/TCR anchors characteristically have a high diversity of targets (“target entropy,” a measure reported by SPLASH) and could be identified on that basis rather than requiring reference mapping. This is expected for clonally diverse receptors and is evident in the clonotypic pattern (each cell expressing only its specific target) seen in heatmaps (Figure 4B).

To showcase SPLASH’s utility for non-model organisms, we ran SPLASH on mouse lemur (Microcebus murinus) samples. Mouse lemurs are primates that diverged from humans 60–75 million years ago and have potential as a model organism. The lemur reference genome is incompletely annotated,
especially at loci such as Ig and TCR. While the human reference does not suffice for alignment-first analysis of mouse lemur transcriptomes, we find that it is a reasonable approximation for interpreting SPLASH outputs; this may generalize to other organisms in which a related, better-curated reference exists.

From Tabula Microcebus, a multi-organ mouse lemur dataset, we analyzed 111 natural killer T (NKT) cells from spleen and, separately, 289 B cells, also from spleen (Tables S2 and S4). In both analyses, the cells came from two different individuals; for this reason, SPLASH also discovered numerous allelic differences between individuals, such as in COX2 (Figure S5A).

Our main focus was on adaptive immune receptors; similar to the human analyses, we found that SPLASH’s lemur anchors in B and NKT cells included Ig C1-set and V-set domains by protein profiling and Ig/TCR gene names by transcript mapping (data not shown). As expected, SPLASH targets for lemur Ig heavy chain are predominantly clonotypically expressed (Figure 4B).

Figure 3. Cell-type-specific expression of paralogs and HLA from single-cell data
(A)–(C) show dot plots, where each dot represents the relative isoform expression in a single cell and the bar marks the average relative expression.

(A) Human MYL12A and MYL12B lie adjacent on chromosome 18, a region syntenic in mammals, chickens, and reptiles. The sequence alignment shows that the two genes are very similar in the coding region and marks the anchor and two targets for individual P2 (P3 has a distinct anchor). Macrophages express relatively more MYL12A and capillary cells more MYL12B, consistent in two individuals.

(B) The HLA-DRB locus occurs as several different haplotypes, all containing DRB1 but differing in paralog (DRB3, DRB4, DRB5, or none; hg38 reference has DRB5). The anchor and its targets (underlined) lie in the 3′ UTRs of DRB1 and DRB4; the two individuals have distinct alleles at both DRB1 and DRB4. Macrophages express mainly DRB1 and capillary cells mainly DRB4.

(C) HLA-DPA1 and HLA-DPB1 overlap in head-to-head arrangement as shown in the UCSC Genome Browser diagram, which also shows the BLAT mapping of the anchor consensuses for DPA1 and DPB1, which lie on opposite strands. This is also depicted in the alignment; the targets are best assigned to opposite strands. An anchor simultaneously reporting on DPA1 and DPB1 was only found for individual P3; its targets show that macrophages exclusively express DPB1, while capillary cells mainly express DPA1.

(D) The polymorphic HLA-B gene contains many SPLASH anchors from T cells, as depicted in the UCSC Browser diagram. The hg38 reference is the B*07:02 allele, whereas this individual matches best to B*08 and B*51 (consensus sequences 1 and 2). We investigated one HLA-B anchor, which lies in exon 2. In the alignment, differences from hg38 in the anchor and lookahead region are in lowercase. In the scatterplot, cells show a wide range of allele expression ratios, with some expressing a single allele. Dashed lines mark a 98% confidence interval for the binomial distribution based on the population average expression (confidence depends on anchor counts); the observed data deviate significantly (binomial p = 1.73E–25), and some cells express almost exclusively one allele.
Figure 4. B and T cell receptor diversity from human and lemur single-cell data

(A) The “transcript mapping” plots show the number of anchors that align to a given gene name, for SPLASH on the y axis and controls on the x axis, with immune receptor genes highlighted in red. For B cells, Ig genes (IgK, kappa; IgL, lambda) predominate among SPLASH anchors but are not found at all in control anchors. For T cells, TCR genes (TRA, alpha; TRB, beta) predominate and are not found in controls. The inset histograms show that immunoglobulin-type “V-set” and “C1-set” are among the top protein domain annotations identified by Pfam on anchor consensuses (for B cells, the top four and bottom four domains are shown; for T cells, all domains are shown). Mobile element activity is suggested by Pfam domains Tnp_22_dsRBD (“L1 transposable element dsRBD-like domain”) in B cells and RVT_1 (“Reverse transcriptase”) in T cells.

(B) Targets associated with Ig/TCR anchors are clonotypically expressed in both human and lemur: heatmaps show that most targets (rows) are expressed only in a single cell (columns). Target sequences are shown as bp color maps (rows are targets, matching the heatmap; columns are bp positions, colored by base) for quick visualization of sequence diversity. Lemur NKT cells show shared TCR usage (see top two rows; the shared target sequence is different in the two individuals).
co-expressing CD3E and KLRB1 (CD161)\textsuperscript{34}; in this cell type, SPLASH also finds shared usage in TCR-beta and TCR-gamma (Figures S5C and SSD).

To test whether SPLASH can find diversity missed by standard methods, we analyzed a subset of 35 lemur B cells for which Ig light-chain variable regions could not be assigned by the program BASIC.\textsuperscript{96} BASIC assigns V(D)J regions based on curated human Ig sequences; although it was able to assign the large majority of lemur B cells, there was a subset for which it failed. We used SPLASH to find evidence of a light-chain variable region in one of the 35 cells (STAR Methods); we were able to reconstruct the full variable region from reads (Data S1). In two cells, there were hits to the surrogate light chain (IGL1/IGL5 or $\lambda5$), which associates with Ig heavy chain when there is not yet a rearranged light chain\textsuperscript{37} (Data S1). This is proof of principle that SPLASH provides insights on data beyond traditional methods. In more recent work, we have built on this capability (J.S. et al., unpublished data).

**SPLASH applied to non-model organisms: Octopus and eelgrass**

To further explore SPLASH’s generality, we applied it to two understudied organisms: octopus and eelgrass. Octopuses have the most complex sensory and nervous systems among invertebrates and are unusual in having high levels of RNA editing.\textsuperscript{38} The marine angiosperm Zostera marina, or eelgrass, is the most widely distributed seagrass, and its adaptation to varying conditions, especially in the face of climate change, is of great interest and is only beginning to be explored at the genomic level.\textsuperscript{39,40}

We focused narrowly on anchors where no more than one of its abundant targets mapped to the respective reference (STAR Methods); that is, where reference-based methods could not have detected variation. Hence, in these analyses, we ignored many interesting findings of SPLASH that are reference consistent.

For octopus, we analyzed an RNA-seq dataset of Octopus bimaculoides\textsuperscript{35} encompassing a variety of tissues from a single individual (N. Bellono, personal communication). We examined several anchors with high effect sizes and BLAST hits to the closely related species Octopus sinensis (STAR Methods; Table S5). An anchor was found in *O. sinensis* myosin-VIla, known as MYO7A in humans; MYO7A mutations cause Usher syndrome, leading to deafness and blindness.\textsuperscript{42,43} Target 1 corresponds to the annotated first exon of *O. sinensis* myosin-VIla, while target 2 represents an alternative first exon (not annotated in either species) expressed only in statocyst tissue (Figure 5A). The *O. bimaculoides* myosin-VIla gene annotation is likely incomplete; it includes neither target, yet both target sequences are found upstream in the reference genome; also, the anchor is missing from the genome (Data S2). The *O. sinensis* myosin-VIla gene is likely misassembled in a different way (Data S2). The statocyst-specific expression of an alternative first exon is intriguing given MYO7A’s association with Usher syndrome and deafness, as the octopus statocyst is a sensory organ for sound and balance,\textsuperscript{43,44} suggesting homologous gene function.

Other SPLASH anchor-targets did not map to the *O. bimaculoides* genome yet did BLAST to 3’ UTRs of *O. sinensis* transcripts, including carboxypeptidase D, Upf2, and netrin receptor/DCC (Figure 6; Table S5). For all three, each target is expressed exclusively in some tissues and not in others. Although *O. bimaculoides* has annotated transcripts for these genes, in all three cases, the 3’ UTR is missing or likely incorrect in the *O. bimaculoides* genome assembly. For two of the genes, the target variation may represent differential expression of alleles: a 13-nt deletion in carboxypeptidase D and a short CAG repeat polymorphism in nonsense-mediated decay gene Upf2 (Figures 6A and 6B). For netrin receptor/DCC, involved in axon guidance and apoptosis, the variation SPLASH detects could be allelic but is also consistent with A-to-I RNA editing (Figure 6C). Our focus here on non-mapping anchor-targets excluded many more examples of regulated variation, including potential RNA editing in numerous octopus genes (data not shown).

We also applied SPLASH to RNA-seq data from eelgrass (*Zostera marina*), collected in two locations: Montpellier, France, (Mediterranean climate) and Rovika, Norway (near-arctic climate), in two seasons (winter and summer) and during day and night.\textsuperscript{40} Considering each anchor with its most abundant target, a large number (14,680, 5.7%) did not align to eelgrass references (Table S6). A high-level view is provided by protein profiling: the top hits were chlorophyll A-B binding protein domain, actin, ubiquitin, and silicon transporter (Figure 5B). Protein BLAST (BLASTP) of some of the translated sequences finds that these have their best hits to a variety of organisms other than eelgrass, notably diatoms. Though a surprise to us, it has long been known that eelgrass is extensively colonized by ephytes,\textsuperscript{45} of which diatoms predominate\textsuperscript{46} and may provide as much as 71%-83% of the primary production by the community.\textsuperscript{47} For the most enriched protein domain, we investigated an anchor with high effect size whose consensus matches “fucoxanthin chlorophyll a/c protein” (FCP) in several diatoms, for example, *Phaeodactylum tricornutum* (95% amino acid identity, 81% nucleotide identity; Figure 7C). Given that the matches are imperfect, the true species of origin may not be in the NCBI database. FCPs function as light-harvesting antennae for photosynthesis.\textsuperscript{48} This anchor has several targets whose abundance varies by location and time of year: target 1 is predominant in France in June; targets 3, 4, and 5, which share the same amino acid sequence, together dominate in France in December; target 6 is predominant in Norway in December (Figure 5C). These targets could represent different diatom species or intra-species allelic variants. The abundance of this anchor (irrespective of target) is lower in night samples (Figure 7C), indicating circadian regulation of this diatom photosynthetic gene. Other anchors mapping to diatoms, such as ferredoxin and high-mobility group box-containing protein, also show targets that segregate by location, France vs. Norway (Figures 7A and 7B).

One anchor and its targets, although mapping to the eelgrass genome, does report on noteworthy variation. It is in the NdhL subunit of chloroplast NADPH dehydrogenase complex (NDH). Of its four most abundant target sequences, three are within exon 3 and are SNP coding variants. The fourth represents retention of the intron following exon 3 (Figure 5D), altering the second transmembrane segment and terminating translation soon after (Data S2). The intron retention variant (target 4) is highest in winter (Figure 5D): for Norway, December (green) vs. June (red) samples.
completely segregate by target 4 expression; the France samples show overlap, but, on average, December (blue) is higher than June (yellow). Figure 5D also illustrates other patterns: Norway samples do not express target 2 (instead they express targets 1 and 3; data not shown); France samples have either a high or a moderate fraction of target 2 (the latter also express target 1; data not shown). NDH is involved in cyclic electron transport, and modulation of NDH function may affect photosynthetic efficiency and oxidative stress under varying light conditions. The above work with octopus and eelgrass is an early foray but shows that SPLASH can discover regulated RNA splicing and isoforms and bring to light allelic variation and communities of associated organisms. SPLASH’s statistics-first and reference-free methodology provides an unbiased approach to discovery, which can be augmented by protein profiling and the use of sequence databases beyond solely genomic references.

DISCUSSION

Genomic analysis today is performed with complex computational workflows that are highly problem specific and reference dependent. Here, we present a unifying statistics-first framework,
A Carboxypeptidase D

O. sinensis

Figure 6. O. bimaculoides 3' UTR anchors show tissue-specific expression

In the heatmaps, the parenthetical numbers are summed anchor counts.

(A) Carboxypeptidase D (CPD). The anchor and targets align to the 3' UTR of the O. sinensis CPD mRNA (XM_029795433.2) but are not in the O. bimaculoides genome assembly. The NCBI Browser screenshot at lower-right shows that the 3' UTR of the O. bimaculoides CPD gene (LOC106880679, Ch.25) is entirely missing from the assembly: immediately after the coding region, a run of Ns begins (red box). Target 2 is identical to O. sinensis except for two mismatches; target 1 has a 12-nt deletion relative to target 2. Target 1 is only expressed in dissociated cells from sucker rims and at a low level in one olfactory organ sample. All other tissues express only target 2.

(B) Upf2 (regulator of nonsense transcripts 2). The alignment of Upf2 mRNAs from O. bimaculoides (XM_014915650.2) and O. sinensis (XM_036513028.1) shows that they diverge just before the stop codon, with unrelated 3' UTRs. Our O. bimaculoides anchor-targets map only to O. sinensis but not to the O. bimaculoides genome assembly. The NCBI Browser screenshot at lower-right shows that the 3' UTR of the O. bimaculoides Upf2 gene (LOC106880679, Ch.25) is entirely missing from the assembly: immediately after the coding region, a run of Ns begins (red box). Target 2 is identical to O. sinensis Upf2 except for two mismatches; target 3 has a 12-nt deletion relative to target 2. Target 1 is only expressed in dissociated cells from sucker rims and at a low level in one olfactory organ sample. All other tissues express only target 2.

(C) Netrin receptor / DCC

In the heatmaps, the parenthetical numbers are summed anchor counts.

In the heatmaps, the parenthetical numbers are summed anchor counts.
SPLASH, which identifies sample-specific sequence variation directly from raw reads using a statistical test.

We provide a snapshot of SPLASH’s discoveries in disparate genomics subfields. When run on SARS-CoV-2 patient samples, without strain metadata or reference genomes, SPLASH finds many anchors capturing strain-defining and emerging mutations. Using reference-free protein domain analysis, SPLASH identifies the spike protein as highly enriched for sequence variation. This points to broad potential for SPLASH in viral and other genomic surveillance.

In single-cell sequencing data, SPLASH is able to identify differential expression between highly similar genes, including myosin light chains MYL12A and -B and several different HLA genes (traditionally difficult to analyze as they are highly polymorphic). SPLASH analysis was conducted in unsupervised mode, yet many of its significant anchors show cell-type regulation between macrophages and capillary endothelial cells. This testifies to the power of SPLASH’s unique statistical approach. When applied to B and T cells of both humans and mouse lemurs, SPLASH automatically identifies antigen receptor genes as exhibiting the most diverse variation. Post facto analysis for lemur was performed using only an approximate genomic reference (human) that diverged from lemur ~60 million years ago.

To examine SPLASH’s ability to find variation not present in reference genomes, we applied it to two diverse organisms: octopus and eelgrass. In octopus, we identified several tissue-regulated isoforms not in the reference, in particular, one in myosin-VIIa that is only expressed in statocyst. In the eelgrass dataset, SPLASH uncovered many sequences from epiphytic diatoms, with variation correlating with geography and season. This highlights the enormous potential in already existing datasets and the need for tools like SPLASH to better explore them.

SPLASH should be of general interest to most genomic analyses. Users can easily run SPLASH on their own samples (FASTQ files): we provide it as a containerized Nextflow pipeline to minimize installation issues; it is lightweight and can be run on a laptop (STAR Methods). The default parameters work well across all tested datasets (and SPLASH is robust to a range of parameters, STAR Methods). SPLASH outputs a list of significant anchors and targets; these results are a large data reduction and distillation of the variation present in the samples, and there are many ways they can be used. If metadata are available, they can be correlated with anchor-targets generated in unsupervised mode; alternatively, one can use metadata to supervise SPLASH analysis. If a reference genome is available, SPLASH can use it to align anchors and targets and provide gene names. SPLASH provides a number of metrics, such as a p value bound, effect size, target entropy, and average target similarity, which can be used to filter the anchor list. Another avenue for analyzing SPLASH’s results is BLAST of anchor-targets or consensus sequences against the NCBI databases and protein domain profiling with databases like Pfam, helpful especially when there is no reference genome or when it is incomplete. Ultimately, users will bring their own domain expertise to bear in deciding how to best utilize SPLASH results.

Even in areas where there are existing pipelines, for example, in differential alternative splicing or antigen receptor identification, SPLASH provides a different approach and may well give additional insights. SPLASH scales to allow discovery to keep pace with the ever-increasing sequence data from the world at large, in particular, microbes and metagenomic communities; recent collaborative work has further increased the computational efficiency of SPLASH.51 SPLASH provides an expansive paradigm and could be applied to a wide range of “omics” modalities, from DNA and protein sequencing to Hi-C and spatial transcriptomics, and more (STAR Methods). The statistical ideas underlying SPLASH are also expansive: anchor-target pairs can be generalized to tensors, and higher-dimensional relations between anchors, targets, and samples can be studied; other functions for splitting and embedding targets and samples can be considered to optimize statistical power.52

In summary, SPLASH shifts from the “reference-first” approach to “statistics first,” performing statistical hypothesis tests on raw sequencing data. By this design, SPLASH is highly computation-efficient. References are valuable for interpretation; however, the filtering of data by reference alignment introduces quantification biases and blind spots. SPLASH promises data-driven biological study with scope and power previously impossible.

**Limitations of the study**
SPLASH can be applied to problems across diverse fields, which are of great current importance (STAR Methods), including those previously discussed. Naturally, some problems are not directly amenable to SPLASH analysis as formulated here. The clearest are cases where quantification of sample-specific RNA or DNA abundance alone is desired (e.g., differential gene expression analysis). Additionally, SPLASH is currently unable to distinguish which biological mechanism underlies the called variation, and work in progress seeks to address this.

**STAR Methods**
Detailed methods are provided in the online version of this paper and include the following:

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O. bimaculoides genome. The alignment also shows the downstream portion of the O. sinensis 3' UTR, where the anchor-targets map. Targets 1 and 2 have six and five tandem CAG repeats, respectively. Target 1 is expressed in dissociated cells from sucker rims and in olfactory organ. The other tissues express target 2. (C) Netrin receptor/DCC. Alignment of the O. bimaculoides genome (gene LOC106883768) and O. sinensis mRNA (XM_03650872.1) shows that the two diverge shortly after the stop codon. The O. bimaculoides gene ends in dinucleotide repeats just before the genome becomes a run of Ns. Our O. bimaculoides anchor-targets map to the O. sinensis netrin receptor 3' UTR but not to the O. bimaculoides genome. The targets differ at a single nucleotide: targets 1 and 2 have G and A, respectively; O. sinensis has a G in this position. If the O. bimaculoides genome encodes A, then target 1 is consistent with A-to-I RNA editing (inosine read as G during reverse transcription). The majority of tissues express target 2 only, while target 1 is only expressed in dissociated cells of sucker rims.
Figure 7. Diatom anchors in eelgrass samples show variation with location/season or day vs. night
(A) HMG (high-mobility group) box domain. The two targets show several nucleotide differences that result in two coding differences. The translation of the consensus sequence has its best two BLASTP matches to HMG box proteins from diatom species, as shown in the inset. Target 1 is found only in Norway/December samples, while target 2 is found only in France/June samples.

(B) Ferredoxin. The two targets show a silent SNP. The translation of the consensus sequence has its best BLASTP matches to ferredoxin from several diatom species; the top two are shown in the inset. Target 1 is found only in France/June samples, while target 2 is found only in Norway/December samples.

(C) Fucoxanthin chlorophyll a/c protein (diatom)

(legend continued on next page)
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.cell.2023.10.028.

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AUTHOR CONTRIBUTIONS
K.C. designed and implemented the pipeline. T.Z.B. designed and analyzed the statistics. I.N.Z. developed the protein domain analysis. P.L.W. spearheaded the later revisions. J.S. designed the statistics and conceptualized and supervised the project. All authors analyzed data and wrote the manuscript.

DECLARATION OF INTERESTS
K.C., T.Z.B., and J.S. are inventors on provisional patents related to this work.

INCLUSION AND DIVERSITY
We support inclusive, diverse, and equitable conduct of research.

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(C) Fucoxanthin chlorophyll a/c protein (FCP). This anchor and its targets are also presented in Figure 5C. At left, the translated consensus sequence has its best protein BLAST matches to several diatom species; two are shown in the inset. The amino acid identity for P. tricornutum is 42/44 (95%). The consensus also BLASTs to the P. tricornutum genome, nucleotide identity 107/132 (81%) (data not shown). At right, histogram shows that total anchor counts for night are ~60% lower than for day, indicating circadian regulation of this gene. All are samples from France in December (where this anchor had both day and night representation).
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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Deposited data      |        |            |
| SARS-CoV-2 France   | Bal et al.⁹ | NCBI: SRP365166 |
| SARS-CoV-2 South Africa | Viana et al.⁸ | NCBI: SRP348159 |
| Rotavirus           | Simsek et al.¹⁷ | NCBI: SRP328899 |
| Human Lung Cell Atlas | Travaglini et al.²⁰ | European Genome-phenome Archive: EGAS000001004344 |
| Tabula Sapiens      | Tabula Sapiens Consortium³⁵ | https://tabula-sapiens-portal.csdbiohub.org/ |
| Tabula Microcebus   | The Tabula Microcebus Consortium et al.³⁴ | https://tabula-microcebus.csdbiohub.org |
| Octopus bimaculoides RNA-seq data | van Giesen et al.⁴¹ | NCBI: SRP327909 |
| Zostera marina RNA-seq data | Jueterbock et al.⁴⁰ | NCBI: SRP327909 |
| Pfam database (Pfam-A) | Mistry et al.⁵³ | https://www.ebi.ac.uk/interpro/download/pfam/ |

Software and algorithms

| SOFTWARE or ALGORITHM | SOURCE | IDENTIFIER |
|-----------------------|--------|------------|
| SPLASH (commit 1b73949) | This study | https://github.com/salzman-lab/SPLASH |
| FASTP v0.23.2 (Installed with bioconda, 2/15/23) | Chen et al.⁵³ | https://github.com/OpenGene/fastp |
| STAR                  | Dobin et al.⁵⁴ | https://github.com/alexdobin/STAR |
| bowtie2               | Langmead et al.⁵₅ | https://bowtie-bio.sourceforge.net/bowtie2/index.shtml |
| bedtools              | Quinlan and Hall⁵⁶ | https://github.com/arq5x/bedtools |
| seqkit (installed with bioconda) | Shen et al.⁵⁷ | https://bioinf.shenwei.me/seqkit/ |
| HMMer3                | Eddy⁵⁸ | http://hmmer.org/ |
| BLAST                 | Camacho et al.⁵⁹ | https://blast.ncbi.nlm.nih.gov/Blast.cgi |

RESOURCE AVAILABILITY

Lead contact
Correspondence and requests for materials should be addressed to the lead contact, Julia Salzman (julia.salzman@stanford.edu).

Materials availability
This study did not generate new unique reagents.

Data and code availability

- This paper analyzes existing, publicly available data. These accession numbers for the datasets are listed in the key resources table.
- All original code has been deposited at Zenodo/Github and is publicly available as of the date of publication. DOIs are listed in the key resources table. Supplemental tables are available for download at https://github.com/salzman-lab/SPLASH/tree/main/cell_data.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

We note that all datasets used are publicly available. We provide relevant details taken from the original papers. Because this work was a study of existing data and because the method can be used on any sample size above two, we did not perform sample size
estimation (all samples in existing data were used). Also, samples were not explicitly allocated to experimental groups, rather the SPLASH procedure described in this manuscript uses random allocation and testing to find significant groupings.

**Human samples**

**South Africa SARS-CoV-2 samples**
Sequencing was done on randomly selected nasopharyngeal and oropharyngeal swab samples from routine diagnostic SARS-CoV-2 PCR testing from public and private laboratories in South Africa and Botswana; requirement for participant consent was waived by the Research Ethics Committees. The large majority of samples were sequenced using Oxford Nanopore, but we only analyzed the samples sequenced using the Illumina COVIDseq assay; the paper does not explain how samples were chosen for Illumina sequencing. The paper does not give a breakdown of samples by age, sex, location, or ethnicity.

**France SARS-CoV-2 samples**
The clinical samples presented in the paper are those that had evidence of co-infection by more than one strain of SARS-CoV-2; they were taken from a broader sequencing effort that included hospitalized patients and health care workers at the university hospital of Lyon (National Reference Center of Respiratory Viruses of Hospices Civils de Lyon (HCL)); and randomly selected positive samples nationwide from all diagnostic laboratories in France (EMERGEN consortium). The study was approved by the HCL ethics committee. We used all the samples in our analysis. Table 1 of the original paper gives some demographic information: among 15 outpatients with Delta/Omicron coinfection, median age was 31.13, 10 were male; 21 outpatients with BA.1/BA.2, median age was 22.36, 6 were male; 13 hospitalized with Delta/Omicron, median age 66.61, 7 were male; 3 hospitalized with BA.1/BA.2, median age 29.03, 2 were male.

**HLCA samples**
Normal lung tissues (approximately 5 cm³) were obtained from uninvolved regions of patients undergoing lobectomy for focal lung tumors; informed consent was obtained. Patient 2 was a 46-year-old male, non-smoker with a right middle lobe (RML) endobronchial carcinoid, who underwent surgical resection of the right upper and middle lobes; two blocks of tissue were selected from mid-bronchial region (medial 2) and periphery (distal 2) of right upper lobe (RUL). Patient 3 was a 51-year-old female, non-smoker with mild adult-onset asthma and a left lower lobe (LLL) endobronchial typical carcinoid, who underwent LLL lobectomy; three tissue blocks were resected from the bronchus (proximal 3), mid-bronchial (medial 2), and periphery (distal 3) of the LLL.

**Tabula Sapiens samples**
Donated tissues were procured in the Northern California through collaboration with the federally mandated organization Donor Network West (DNW, San Ramon, CA, USA); the research protocol was approved by the relevant boards of DNW and Stanford University. Donor TSP1 was a 59-year-old female with BMI of 23 and a history of stroke. She was found down with slurred speech. Donor TSP2 was a 61-year-old female with BMI of 41 and a history of deep vein thrombosis, hypertension, hypersensitive lung disease, type 2 diabetes, an infected knee replacement, and recent bout of the flu. She reported being short of breath and later went into respiratory arrest.

**Mouse lemur samples**
*Microcebus murinus* gray mouse lemurs originated from the closed captive breeding colony at the Muséum National d’Histoire Naturelle in Brunoy, France, and transferred to the University of Texas (Austin) and then Stanford University. Mouse lemurs were housed indoors in an AAALAC-accredited facility in a temperature (24°C) and light-controlled environment (daily 14:10 h and 10:14 h light: dark alternating every 6 months) with perches and nest boxes, and were fed fresh fruits and vegetables, crushed primate chow plus live insect larvae. Animals in declining health that did not respond to standard therapy were euthanized by pentobarbital overdose under isoflurane anesthesia. Organs and tissues were removed and divided by a veterinary pathologist. We used data from two individuals: L2, 10 year old female; and L4, 11 year old male.

**Octopus samples**
Adult female California two-spot octopuses (*Octopus bimaculoides*) were wild-caught (Aquatic Research Consultants, San Pedro, CA), fed daily with fiddler crabs (*Uca pugnax*, Northeast Brine Shrimp, Oak Hill, FL), and kept on a 12h light/dark cycle in natural sea water. Sensory cells were isolated from suckers following tissue extraction and sucker epithelium dissection from animals that were sedated using stepwise increases in ethanol (ending at 3%). Animal protocols were approved by the Harvard University Animal Care and Use Committee.

**Eelgrass (Zostera marina) samples**
Eelgrass shoots were sampled in Norway (Røvik, 67°16’06.2” N, 15°15’38.4” E) and France (Sète, Thau Lagoon, 43°25’08.0” N, 3°40’03.9” E). The youngest two shoots of each of six plants were collected at each site at noon and the following midnight around summer solstice (June 21) and winter solstice (December 21) 2017.

**METHOD DETAILS**

**SPLASH overview**
Full details of SPLASH usage and outputs can be found at [https://github.com/salzman-lab/SPLASH](https://github.com/salzman-lab/SPLASH). Briefly, it takes as input a set of FASTQ sequencing data files (one per sample). SPLASH has several tunable parameters (anchor and target length, lookahead,
minimum count thresholds, and more). SPLASH’s default settings work well in practice, and SPLASH’s inference is robust to these choices (see below). The standard output includes a table of anchors, targets, p values, etc., a table of “consensus” sequences (see below), and a table of “element annotations” (see below). SPLASH can also perform alignments with bowtie2 and STAR, to generate “genome annotations” and splice junction annotations (see below).

The code used in this work is available as a fully-containerized Nextflow pipeline at https://github.com/salzmanlab/SPLASH, commit 1b73949. The GitHub repository also contains the sample sheets for all analyses, including individual sample SRA accession numbers; as well as scripts for supplemental analysis. See quantification and statistical analysis, below, for explanation of SPLASH p value and effect size computations.

**SPLASH anchor preprocessing and parameter choices**

Anchors and targets are defined as sequences of length \( k \) (k-mers) positioned at an offset \( R \) apart. By default, \( R = \text{max}(0, (L - 2^k)/2) \) where \( L \) is the length of the first read processed in the dataset, and \( R \) is rounded to the nearest integer (e.g., if \( L = 100 \) and \( k = 27 \), then \( R = 23 \)). For a fixed number of anchor-target pairs, under alternatives such as differential exon skipping, larger choices of \( R \) have provably higher power than smaller choices, following the style of analysis in Salzman et al.\(^61\) \( k = 27 \) is typically long enough to be assigned a unique position in a genome while having a low probability of containing a sequencing error. Anchor sequences can be extracted as adjacent, disjoint sequences or as tiled sequences that begin at a fixed step size, to reduce computational burden. For this manuscript, SPLASH was run with default parameters: with 1 M reads per FASTQ file, anchor sequences tiled by 5 bp, and \( k = 27 \). For HLCA datasets, both read 1 and read 2 were used; for other datasets, only read 1. Extracted anchor and target sequences are then counted for each sample with the UNIX command, ‘sort | uniq -c’, and anchor-target counts are then collected across all samples for reannotation by the anchor sequence. This stratification step allows for user control over parallelization. To reduce the number of hypotheses tested and required to correct for, we discard anchors that have only one unique target, anchors that appear in only 1 sample, and (anchor, sample) pairs that have fewer than 6 counts. Then, we retain only anchors having more than 30 total counts after the above thresholds were applied. This approach efficiently constructs sample by target counts tables for each anchor.

SPLASH is robust to parameter choices. We give examples of how choices of \( k, R, \) and tiling length impact results in France SARS-CoV-2 data as follows, showing that SPLASH yields similar results for a range of parameter choices. Default parameters shown in bold: we tested \( k = [25, 27, 30] \); Tile = \([3, 5, 7]\); Lookahead = \([0, 15, 23]\). For \( k = 25 \), 94.4% of anchors with default parameters contain at least one of the K = 25 anchors as a substring. For \( k = 30 \), 93.8% of anchors with \( k = 30 \) contain at least one of the anchors with default parameters as a substring. For tile size of 3, 85% of the anchors from the default run can be found in the significant anchors of tile size of 3. For tile size of 7, 85% of the anchors from the default run can be found in the significant anchors of tile size of 5. For lookahead distance of 0, 37% of the anchors from the default run can be found in the significant anchors of tile size of 3; for lookahead distance of 15, 76% of the anchors from the default run can be found in the significant anchors. Overall, as tile size decreases, anchor calls increase (4715, 5522, 7891 for \([7, 5, 3]\) respectively). As \( k \) varies, anchor calls stay essentially the same (5875, 5522, and 5958 for \( k = [25, 27, 30] \) respectively). Finally, for lookahead distance, the total number of calls decrease as lookahead distance increases (13239, 8295, 5522 for \( R = [0, 15, 23] \) respectively).

**Consensus sequences**

For each significant anchor, a per sample consensus sequence is built for the sequence downstream of the anchor. A separate consensus is built for each sample by aggregating all reads from this sample that contain the given anchor. Then, SPLASH constructs the consensus as the plurality vote of all these reads; concretely, the consensus at base pair \( i \) is the plurality vote of all reads that contain the anchor, \( i \) base pairs after the anchor appears in the read (a read does not vote for consensus base \( i \) if it has terminated within \( i \) base pairs after the anchor appeared). The consensus base as well as the fraction agreement with this base among the reads is recorded. Some empirical behavior of consistencys is shown in Figure S4.

The consensus sequences can be used for splice site discovery as well as other applications, such as identifying point mutations and highly diversifying sequences, e.g., V(D)J rearrangements. The statistical properties of consensus building make it an appealing candidate for use in short read sequencing,\(^62\) and may have information theoretic justification in de novo assembly.\(^63\)

To provide intuition regarding the error correcting capabilities of the consensus, consider a simple probabilistic model where our reads from a sample all come from the same underlying sequence. In this case, under the substitution only error model, we have that the probability that our consensus for \( n \) reads makes a mistake at a given location \( i \) under independent sequencing error rate \( \varepsilon \) (substitution only) is at most

\[
\Pr(\text{error at basepair } i) \leq \sum_{k \geq n/2} \binom{n}{k} \varepsilon^k (1 - \varepsilon)^{n-k} \leq \frac{n}{n/2} \varepsilon^{n/2}.
\]

We can see that even for \( n = 10 \), this probability is less than \( 1.3 \times 10^{-7} \) for a given base pair, which we can union-bound over the length of the consensus to yield a vanishingly small probability of error. Thus, for a properly aligned read, if a base pair differs between the consensus and reference it is almost certainly an SNP.

**Element annotations**

To identify false positive sequences or contextualize mobile genetic elements, anchors and targets are aligned with bowtie2 to a set of indices, corresponding to databases of sequencing artifacts, transposable elements, and mobile genetic elements.\(^64\) In these
alignments, using bowtie2, the best hit is reported, relative to an order of priority. The references used are: UniVec, Illumina adapters, grass carp (Ctenopharyngodon idella genome, GCA_019924925.6), Escherichia phage phiX174, Pfam, Dfam, TrnCentral, ACtME, Iceberg, CRISPR direct repeats, ITSoneDB, ITS2, and also the reference genome of interest for the study. (Grass carp was used as a control as it contains many artifactual Illumina adapters.) To perform these annotations, bowtie2 indices were built from the respective reference FASTAs, using bowtie2-build with default parameters. Anchors and targets were then aligned to each index, using bowtie2-align with default parameters. For each sequence, we report the alignment to the reference and the position of that alignment for each reference in the prespecified set. Anchors and targets, and their respective element annotations, are reported in the element annotation summary files.

**Genome annotations**

Anchor, target, and consensus sequences can be aligned by SPLASH to reference genomes and transcriptomes, to provide information about the location of sequences relative to genomic elements. All alignments reported are run in two modes in parallel: bowtie2 end-to-end mode (the bowtie2 default parameters) and bowtie2 local mode (`-local`, in addition to the bowtie2 default parameters). To report alignments to the transcriptome, the sequences are aligned to the reference transcriptome with bowtie2, with `-k 1`, in addition to the above parameters, to report a maximum of one alignment per sequence. If there is a transcriptome alignment, we report the alignment to the reference and the MAPQ score of the alignment. To report alignments to the genome, the sequences are aligned to the reference genome, with the same parameters above. If there is a genome alignment, we report the alignment to the reference, the strand of the alignment, and the alignment MAPQ score.

**Splice junction calls**

To identify exon coordinates for reporting annotations in this manuscript, consensus sequences are mapped with STAR aligner (default settings). Gapped alignments are extracted and their coordinates are annotated with known splice junction coordinates using ‘bedtools bamtobed –split’; each resulting contiguously mapping segment is called a “called exon”. From each consensus sequence, called exons are generated as start and end sites of each contiguously mapped sequence in the spliced alignment. These ‘called exons’ are then stratified as start or end sites and end sites. Note that the extremal positions of all called exons would not be expected to coincide with a splice boundary; “called exon” boundaries would coincide with an exon boundary if they are completely internal to the set of called exon coordinates. Each start and end site of each called exon is intersected with an annotation file of known exon coordinates; it receives a value of 0 if the site is annotated, and 1 if it is annotated as alternative. The original consensus sequence and the reported alignment of the consensus sequence are also reported. Gene names for each consensus are assigned by bedtools intersect with gene annotations (hg38 RefSeq for human data by default), possibly resulting in multiple gene names per consensus.

**SPLASH protein domain profiles**

Custom scripts were used to generate protein domain profiles. For each set of enriched anchors, homology-based annotation was attempted against an annotated protein database, Pfam. For each dataset, up to 1000 of the most significant anchors (q-value < 0.01) were retained for the following analysis: we first generated a substring of each downstream consensus by appending each consensus nucleotide assuming both conditions were met: a minimum observation count of 10 and a minimum agreement fraction of 0.8, until whichever metric first exhibited two consecutive failures at which point no further nucleotide was added. A limit of 1000 anchors was used due to computational constraints from HMMer3 (see below). Anchors that did not have any consensus nucleotides appended were kept as is. An extended anchor was generated for each experiment in which an anchor was found. Each extended anchor was then stored in a final concatenated multi FASTA file with unique seqID headers for each experiment’s extended anchors.

To assess these extended anchors for protein homology, this concatenated FASTA file was then translated in all six frames with the standard translation table using seqkit prior to using hmmssearch from the HMMer3 package to assess resulting amino acid sequences against the Pfam35 profile Hidden Markov Model (pHMM) database. The resulting ‘raw’.tblout outputs were then processed, keeping the best hit (based on E-value) per each initial anchor, and any hits with an E-value better than 0.01 were parsed into an ‘.nomad.Pfam’ (or ‘.control.Pfam’) file used for subsequent plotting.

All hits to the Pfam database were then binned at different E-value orders of magnitude. In each case, control assessments were performed by repeating the extension and homology searches against an equivalent number of control anchors (see below). Protein domains are ranked in the plot by the difference between SPLASH anchor hits and control anchor hits (for hits with E-value ≤ 1e-02).

The number of matched anchors used for SPLASH and control analysis per dataset were as follows: 201 high effect size (0.5) anchors in SARS-CoV-2 from South Africa, 252 high effect size (0.5) anchors in SARS-CoV-2 from France; 1000 anchors (no effect size filter) were used for rotavirus, human T cells, human B cells, Microcebus natural killer T cells, and Microcebus B cells. We note that while the number of input anchors for SPLASH and control sets are matched, it is possible to have more control protein domains in the resulting barplots, as only high E-value hits to Pfam are reported in the visualizations. Domain profiling summaries are in Data S3.

A hypergeometric test was used to give p values for protein domain analysis. For a given domain, we construct the 2 × 2 contingency table, where the first row is the number of SPLASH hits for this domain, followed by the total number of SPLASH hits not in this domain. The second row is the mirror of this for control, where the first entry is the number of control hits for this domain, followed by the total number of control hits not in this domain. A one-sided p value is computed using Fisher’s exact test, which is identically a
SARS-CoV-2 data were downloaded from the NCBI: France\(^8\) (SRP365166) and South Africa\(^8\) (SRP348159). Sample metadata for the SARS-CoV-2 analysis corresponding to additional targets based on biological relevance. All samples with fewer than 5 counts are discarded. For clarity of presentation, we include or remove rows than 5 times, have less than 5% of the total counts of that anchor, and retain at most the top 10 targets, while ensuring that at least 2 targets are plotted. Then, all samples with fewer than 5 counts are discarded. For clarity of presentation, we include or remove rows corresponding to additional targets based on biological relevance.

Control analyses
To construct control anchor lists based on abundance, we considered all anchors input to SPLASH and counted their abundance, collapsing counts across targets. That is, an anchor receives a count determined by the number of times it appears at an offset of 5 in the read up to position \(R - \max(0,R/2-2*k)\) where \(R\) is the length of the read, summed over all targets. The 1000 most abundant anchors were output as the control set. For analysis comparing control to SPLASH anchors, \(\min(\text{SPLASH anchor list}, 1000)\) most abundant anchors from the control set were used and the same number of SPLASH anchors were used, sorted by p value.

Generation of contingency table heatmaps
To plot the anchor-target heatmaps, we exclude targets with low counts. Concretely, we by default filter out targets that occur fewer than 5 times, have less than 5% of the total counts of that anchor, and retain at most the top 10 targets, while ensuring that at least 2 targets are plotted. Then, all samples with fewer than 5 counts are discarded. For clarity of presentation, we include or remove rows corresponding to additional targets based on biological relevance.

SARS-CoV-2 analysis
SARS-CoV-2 data were downloaded from the NCBI: France\(^8\) (SRP365166) and South Africa\(^8\) (SRP348159). Sample metadata for the France dataset was provided by the authors via personal communication, and consists of their calls of the primary and secondary infecting strains for each patient sample. We note that sample ‘WTA-022002271301_S1’ appeared to be mislabeled, appearing in the metadata file but not in the NCBI sample list. Conversely, the sample ‘PI924-022002271301_S1472’ appears in the NCBI sample list, but not in the metadata file. Thus, we associate these labels to each other, to obtain metadata labeling for all 106 samples. We do not have information regarding which samples are replicates. We provide the NCBI sample list and the strain metadata file in Table S1.

The SARS-CoV-2 datasets used in this manuscript were analyzed with SPLASH’s unsupervised mode (no sample metadata provided). To identify high effect size anchors, a threshold of ‘effect_size_randCjs’ > 0.5 was used (Table S2).

For the purposes of strain-defining mutation analysis, we manually constructed “archetype” genome sequences for variant strains Delta, Omicron BA.1, and Omicron BA.2 by editing the Original (Wuhan) reference NC_045512.2 to contain all (and only) the defining mutations specified at CoVariants.org\(^{10}\); these are provided in Table S1.

To determine what SPLASH calls (and control anchors) were strain defining we perform the following. To generate SPLASH’s calls, we filter for anchors that are significant (with a BY corrected p value less than 0.05) and have large effect size (>0.5), yielding a list of \(N\) SPLASH-called anchors. Control anchors are generated by taking the \(N\) anchors with the highest counts. For each of these anchors we construct their target \(\times\) sample contingency table, first filtering out all anchors with fewer than 30 counts, only 1 unique target, or only 1 unique sample, and filtering out all samples with 5 or fewer counts. Then, we discard all targets that constitute less than 5% of the remaining counts for that anchor. The remaining anchors and targets are then bowtie aligned to an index comprised of the Original, Delta, Omicron BA.1, and Omicron BA.2 archetype genomes. For this alignment, options ‘-a -v 0’ were used. Then, for each set of anchors (SPLASH calls, and controls), the list is filtered to only anchors that align perfectly to at least one of the reference assemblies, further requiring that each anchor have at least one target that aligns perfectly to a reference assembly. Then for each anchor, we declare it to be strain defining if, for any of the reference assemblies, it has at least one target that maps to it and one target that does not.

Identifying cell-type-specific isoforms in single-cell data (lung macrophages and capillary cells)
The human lung scRNA-seq data used here (HLCA SS2\(^{20}\)) is accessible through the European Genome-phenome Archive (accession number: EGAS00001004344); FASTQ files from donor 1 (P2) and donor 2 (P3) generated with the Smart-seq2 protocol were used. In the analysis of HLCA SS2 data, we utilize “isoform detection conditions” for alternative isoform detection. These conditions select for (anchor, target) pairs that map exclusively to the human genome, anchors with at least one split-mapping consensus sequence, \(mu_{-}lev > 5\), and \(M > 100\); \(mu_{-}lev\) is the average target distance from the most abundant target as measured by Levenshtein distance, \(M\) is the total number of counts in the anchor’s contingency table. To identify anchors and targets that map exclusively to the human genome, we included anchors and targets that had exactly one element annotation, where that one element annotation must be gRch38_1kgmaj. To identify anchors with at least one split-mapping consensus, we selected anchors that had at least one consensus sequence with at least 2 called exons. The conditions on Levenshtein distance, designed to require significant across-target sequence variation, significantly reduced anchors analyzed (excluding many SNP-like effects). We further restricted to anchors with \(M > 100\), to account for the lower numbers in macrophage cells; note that the user can choose to use a lower \(M\) requirement, based on input data. These isoform detection parameters were used to identify the SS2 examples discussed in this manuscript. For HLA discussion, gene names were called using consensus\_gene\_mode.
While here we focus on anchors that have aligned to the human genome, we note in passing that SPLASH makes many predictions of cell-type-specific RNA expression that include sequences that map to repetitive elements or do not map to the human reference: for individual P2 (respectively P3), 53% (61%), of 4010 (4603) anchors map to the human genome and no other reference; 35% (30%) map to both the Rfam and human genome; 6% (7%) have no map to any reference used for annotation which includes repetitive and mobile elements. As an example, 9 and 18 such anchors (individual P2 and P3, respectively) BLAST to MHC alleles in the NCBI database.

**Immune single-cell analysis**

To study human B and T cells, we utilize Tabula Sapiens data (Smart-seq2), downloaded from https://tabula-sapiens-portal.ds.czbiohub.org; B cells were used from donor 1 (TSIP1) and CD4+ T cells from donor 2 (TSP2). Mouse lemur single-cell RNA-seq data used in this study was generated as part of the Tabula Microcebus consortium; the FASTQ files were downloaded from https://tabula-microcebus.ds.czbiohub.org. B cells and natural killer T cells were analyzed separately; both were from spleen and were a mixture of individuals L2 and L4. To determine the most frequent transcriptome annotation for a dataset, all significant anchors were mapped to the human transcriptome (GRC38, Gencode) with bowtie2, using default parameters and `-k 1` to report at most one alignment per anchor (Table S4). Then, the bowtie2 transcript hits are aggregated by counting over anchors. The transcript hits with the highest counts over all anchors were reported. Protein domain profiling was performed as described above.

**SPLASH comparison to BASIC analysis in lemur spleen B cells**

As part of the Tabula Microcebus consortium, mouse lemur B cells were annotated with BASIC to identify Ig variable domains. However, BASIC was unable to identify the light chain variable domain in 35 cells. We used a simple approach to see if SPLASH could identify variable domains in these uncalled B cells. We checked the SPLASH genome annotations for these cells for anchors mapping to human “IgL” or “IgK” genes; there were only five such anchors, all to IGL, and these were found in only eight cells. For those eight cells, we retrieved the SPLASH consensus sequences for these anchors, which ranged from 2 to 5 per cell. Where consensus sequences for a cell overlapped, one was chosen, and these were submitted to BLAST against the nr/nt database. Many hits were to “immunoglobulin lambda-like polypeptide” 1 or 5 (IGLL), surrogate light chain genes that contain sequence similar to lambda J and C regions (as well as a unique N-terminal region) and so could mimic alignment to a true lambda variable domain. Therefore, BLAST alignments were checked to see whether the match could be assigned to V, J, C, or IGLL-unique regions. 4 cells matched C-region, 2 matched IGLL-unique region, and 2 had sequence beyond J-region (presumed V-region). For the latter two, we attempted to extend the consensus further into the V-region by ‘grep’ in raw reads; one could not be extended as it only had adaptors adjacent to its J-region sequence. The other consensus was extended through the full V-region, and its sequence is given in Data S1, along with the IGLL-unique matches.

**SPLASH for Zostera marina (eelgrass) and Octopus bimaculoides**

Data was downloaded from SRP327909 (eelgrass), and SRP278619 (Octopus), using nf-core fetchngs run in default mode and preprocessed with fastp run in default mode to mitigate false positive calls due to adapter concatenation. An updated version of SPLASH (R-SPLASH v0.3.9, commit 5dafdc8) was run with gap length = 0, anchor_unique_targets_threshold = 1, anchor_count_threshold = 50, anchor_samples_threshold = 1, anchor_sample_counts_threshold = 5, and excluding anchor-targets containing poly A/C/G/T run of length 8. 500 pairs of random c and f were chosen. fastp v0.3.9 was installed on 2/23/23 using bioconda.

The top 10 anchor-targets for each anchor were selected from the SPLASH calls if they had homopolymer length ≤ 5, effect size > 0.1, and corrected p value < 0.01. Element annotations were run using SPLASH commit ID 728066b; anchor-targets mapping to UniVec, Illumina adapters, SARS-CoV-2, or grass carp were removed. Anchor-targets were aligned with STAR 2.7.5 to a reference index generated from either the O. bimaculoides reference genome (NC_068981.1) and transcriptome; or the Z. marina genome (v3.1, https://phytozome-next.jgi.doe.gov/info/Zmarina_v3_1) and mitochondrial and chloroplast genomes (NC_035345.1 and NC_036014.1, respectively).

For protein domain profiling, anchor-targets with no element annotation were in silico translated in all six frames and submitted to HMMer search of the Pfam database. For Figure 5B (eelgrass), anchors were ordered by descending number of observations; the top 200,000 were concatenated with their targets and submitted to element annotations, and unannotated anchor-targets were submitted to Pfam; these anchor-targets are defined as controls. For each anchor, we retain its best Pfam hit and full sequence E-value. Then for each Pfam domain, we tally (separately for SPLASH and control) the number of anchors that hit it with a full-sequence E-value ≤ 1e-02 and ≤ 1e-06.

For BLAST analysis, we selected anchors with no more than 1 target mapping with either STAR to the reference genome or Bowtie2 to human genome, thus cases where no sample-specific variation would be detected if a reference genome were used. In eelgrass, targets were selected if their fraction exceeded 0.5 and their anchor’s effect size exceeded 0.9; in octopus, targets were selected if their fraction exceeded 0.3 and if their anchor’s effect size exceeded 0.8. The 1808 anchor-target pairs satisfying these criteria in octopus were submitted to BLAST with parameters:

- `db nt -evalue 0.1 -task blastn -dust no -word_size 24 -reward 1 -penalty -3 -max_target_seqs 4`

BLAST hits were merged into SPLASH output, with an indicator variable for whether the sequence was queried. For octopus anchor-targets, 1061/1808 had a BLAST hit (max E-value 0.028). There were 288 hits to octopus, of which 281 were annotated as from
O. sinensis and 7 from O. bimaculoides. Selected sequences mapping to O. sinensis were further analyzed. For eelgrass, 1606/4081 had a BLAST hit (max E-value 0.028).

SPLASH output, merged with Pfam and BLAST analyses, are in Tables S5 and S6 (octopus and eelgrass, respectively).

**SPLASH can detect myriad genomic events**

In this work we focused our experimental results on identifying changes in viral strains and specific examples of RNA-seq analysis. SPLASH's probabilistic formulation extends much further however, and subsumes a broad range of problems. Many other tasks, some described below, can also be framed under this unifying probabilistic formulation. Thus, SPLASH provides an efficient and general solution to disparate problems in genomics. We outline examples of SPLASH's predicted applications in various biological contexts, highlighting the anchors that would be flagged as significant.

- RNA splicing, even if not alternative or regulated, can be detected by comparing DNA-seq and RNA-seq
  - Examples of predicted significant anchors: sequences upstream of spliced or edited sequences including circular, linear, or gene fusions
- RNA editing can be detected by comparing RNA-seq and DNA-seq
  - Examples of predicted significant anchors: sequences preceding edited sites
- Liquid biopsy – reference free detection of SNPs, centromeric and telomeric expansions with mutations
  - Examples of predicted significant anchors: sequences in telomeres (resp. centromeres) preceding telomeric (resp. centromeric) sequence variants or chromosomal ends (telomeres) in cancer-specific chromosomal fragments
- Detecting MHC allelic diversity
  - Examples of predicted significant anchors: sequences flanking MHC allelic variants
- Detecting disease-specific or person-specific mutations and structural variation in DNA
  - Examples of predicted significant anchors: sequences preceding structural variants or mutations
- Cancer genomics e.g., BCR-ABL fusions
  - Examples of predicted significant anchors: sequences preceding fusion breakpoints
- Transposon or retrotransposon insertions or mobile DNA/RNA
  - Examples of predicted significant anchors: (retro)transposon arms or boundaries of mobile elements
- Adaptation
  - Examples of predicted significant anchors: sequences flanking regions of DNA with time-dependent variation
- Novel viruses and bacteria; emerging resistance to human immunity or drugs
  - Examples of predicted significant anchors: sequences flanking rapidly evolving or recombined RNA/DNA
- Alternative 3’ UTR use
  - Examples of predicted significant anchors: 3’ sequences with targets including both the poly(A) or poly(U), or adapters in cases of libraries prepared by adapter ligation versus downstream transcript sequence
- Hi-C or any proximity ligation
  - Examples of predicted significant anchors: for Hi-C, DNA sequences with differential proximity to genomic loci as a function of sample; similarly, for other proximity ligation anchors would be predicted when the represented element has differential localization with other elements
- Finding combinatorially controlled genes e.g., VDJ
  - Examples of predicted significant anchors: sequences in the constant, J, D, or V domains

**SPLASH can use alternative anchor, target and consensus construction**

SPLASH can function on any biological sequence and does not need anchor-target pairs to take the form of gapped k-mers, and can take very general forms. For example, one could consider schemes that respect triplet codons: \[X_1X_2Y_1][X_3X_4Y_2][X_5X_6Y_3] \ldots \] where \(X_i\) are bases in the anchor and \(Y_i\) are bases in the target, this would focus specifically on variation in the wobble position, the fastest to diverge; similar schemes might be appropriate for mechanisms with known patterns of diversity, such as diversity generating retro-elements.\(^7^9\) \(X\) and \(Y\) could also be amino acid sequences or other discrete variables considered in molecular biology. Much more general forms of anchor-target pairs (or tensors) can be defined and analyzed, including using univariate or multivariate functions on targets or sample identity. SPLASH can also be further developed to analyze higher dimensional relationships between anchors, where statistical inference can be performed on tensors across anchors, targets, and samples. Similarly, these functions can be optimized under natural maximization criterion, which is the subject of concurrent work.\(^5^2\) These functions can also be generalized to yield new statistics, optimizing power against different alternatives.

**Computational benchmarking for SPLASH**

SPLASH is computationally much more efficient than other approaches, due to its use of k-mers rather than reference alignment, and its closed-form statistics obviating compute-intensive significance testing. SPLASH is implemented as a fully containerized and parallelized workflow that requires only the FASTQ read files and no parameter tuning by the user. We ran SPLASH on a 2015 Intel laptop.
with an Intel Core i7-6500U CPU @ 2.50GHz processor, generating significance calls for single cell RNA-seq totaling over 10 million
reads in only 1 h 45 min. When performed on a compute cluster, the same analysis is completed in an average of 22.8 min with 750
MB of memory for 10 million reads.

Because code was run on a server with dynamic memory, we report summary statistics as follows. For the steps parallelized by
FASTQ file, such as anchor and target retrieval, total time for dataset run, as reported by Nextflow, was parsed per cell. Thus, the
average time per cell is reported. For the steps parallelized by 64 files (q value calculations), total extracted times were summed
and divided by number of cells. For steps that consisted of aggregating files, total run time was divided by number of cells. Thus,
the total time and memory should be multiplied by the total number of cells to achieve an estimate of the pipeline time for this dataset.

**Laptop specs**

An Intel Core i7-6500U CPU @ 2.50GHz (launched in 2015)
2 cores, total of 4 threads, 3 of which SPLASH was allowed to use.
8 GB DDR3 RAM.
SODIMM DDR3 Synchronous 1600 MHz (0.6 ns)

**Laptop analysis dataset**

Ten B and T cells from donor 2 blood sequenced by Smart-Seq2 were used for the laptop benchmarking. These files totaled
43,870,027 reads, averaging 4.3M reads per cell. The fastq files for the Tabula Sapiens data were downloaded from https://
tabula-sapiens-portal.ds.czbiohub.org/.

Files used:

TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A13_S73_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A18_S78_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A19_S79_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A21_S81_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A3_S63_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A5_S65_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A6_S66_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A8_S68_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_A9_S69_R1_001.fastq.gz
TSP2_Blood_NA_SS2_B114581_B133053_Lymphocytes_B10_S94_R1_001.fastq.gz

**Anchor and target sequences, with q values and binomial p value bounds**

Targets are numbered by decreasing abundance, unless otherwise stated.

q values are the BY-corrected p value bounds output by SPLASH, as detailed in quantification and statistical analysis, below.
Binomial p value bound calculations are described below, and are with respect to target 1, unless otherwise stated.

**SARS-CoV-2 mutation K417N (Figure 2A)**

q value: 9.4e-05
binomial p value: 6.4e-07.
>anchor
ATTCAATTGTATTTAGAGGCTGATGAAG
>target_1_Delta
ACTGGAAGATAGCTGATTATAATTAT
>target_2_K417N_Omicron
ACTGGAATATTGCTGATTATAATTAT

**SARS-CoV-2 mutations V213G, NL211I, R214REPE (Figure 2B)**

q value: 8.3e-08
binomial p value: 1e-13.
>anchor
TTTAAAGATATATGATGTTATTTTTAAA
>target_1_Delta
TAAATTAGGCTGATCTCCCTCAGGG
>target_2_V213G_BA.2
TAAATTAGGCTGATCTCCCTCAGGG
>target_3_NL211I-R214REPE_BA.1
TATGGGCTGATCAGAGATCTCCC

**SARS-CoV-2 mutations P681R, N679K, P681H (Figure 2C)**

q value: 1.2e-04
binomial p value: 4.9e-12.
(reverse-complements are shown in Figure 1C)

>anchor
GTGACATAGTGTAGGCAATGATGGATT
>target_1_P681R_Delta (abundance order = 1)
CGACGAGAATTAGTCTGAGTCTGAT
>target_2_P681R-Q677H (abundance order = 3)
CGACGAGAATTAGTCTGAGTCTGAT
>target_3_P681R-Q677H (abundance order = 4)
CGACGAGAATTAGTCTGAGTCTGAT
>target_4_N679K-P681H_Omicron (abundance order = 2)
CGATGAGACTTAGTCTGAGTCTGAT

**MYL12A/MYL12B (Figures 3A and S4B)**
P2 q value: 2.5e-08.
P2 binomial p value: 9.9e-37 (with respect to target 2)
P3 q value: 2.3E-42.
P3 binomial p value: 2.2e-45 (with respect to target 1)

>P2_anchor
AAGAGCCCTTCAACATGATGGAT
>P2_target_2_MYL12A
TTCCATGGGAAGAATCCAACTGAT
>P2_target_1_MYL12B
TTCTTGGAAGAATCCAACTGAT
>P2_consensus_MYL12A_macrophage
ACAGAGATGGTTTCATCGACAAGGAAGATTTGCATGAGTCTGCTCTCATTGGGAAGAATCCAACTGATGAGGCTCCAGGCC
>P2_consensus_MYL12B_capillary
ACAGAGATGGCTTCATCGACAAGGAAGATTTGCATGAGTCTGCTCTCATTGGGAAGAATCCAACTGATGAGGCCCCAGGCC

**HLA-DRB1/HLA-DRB4 (Figure 3B)**
P2 q value: 4.0e-10.
P2 binomial p value: 9.9e-37
P3 q value: 1.2e-4.
P3 binomial p value: 1.6e-08.

>anchor
GGAAGCCACAAGGGAGGACATTCTTG
>target_1_HLA-DRB1
GTGGAAGAATAACTGCCAAGCAGGAA
>target_2_HLA-DRB4
GGAAGAATAAGAGCCAAGTGGGAAAGC
>p3_consensus_HLA-DRB1_macrophage
GGAAGCCACAAGGGAGGACATTCTTG
>CAGTTGCCGAACCAGTAGCAACCAGT
>target_2_HLA-DRB4_capillary
GGAAGAATAAGAGCCAAGTGGGAAAGC
>target_1_HLA-DRB1_capillary
GGAAGCCACAAGGGAGGACATTCTTG
>CAGTTGCCGAACCAGTAGCAACCAGT

HIGHLIGHTS

- Both MYL12A and MYL12B are expressed in the macrophage and capillary subtypes.

The expression levels of MYL12A and MYL12B are significantly different between macrophages and capillaries, suggesting that their expression may play a role in the function of these cell types.
CAGAGTGCTGAAACAGCATACCTCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

> P3anchor (same sequence as P2anchor)
GGAAGGCTCTCCTCCTTCCTTCCTTCTTTGG

> P3_target_1_DRB1
AGGTGTCAAGAAGCCTCTCTTGAGG

> P3_target_2_DRB1
CCTGGTCCCTGACAAAGCTCTTGGAA

> P3_consensus_DRB1_macrophage
CAGTTGCTGAAACAGCATACCTCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

GGGAGAAAGGTTCTTCT

> P3_consensus_DRB4_capillary
GGAAGGCTCTCCTCCTTCCTTCCTTCTTTGG

CAGAGTGCTGAAACAGCATACCTCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

HLA-DPA1/HLA-DPB1 (Figures 3C and S4C)
P3 q value: 7.9e-22.
P3 binomial p value: 9.15e-18.
(anchor as given here is sense strand for DPA1, antisense strand for DPB1)

> P3anchor
AGATGATCTCTCCTGAAAGGCTCTTCT

> P3_target_1_DPA1
TGCCGTCCCTGGAAAGTGAAATCCCA

> P3_target_2_DPB1
TGCCGTCCCTGGAAAGTGTAATTCTCT

> P3 consensus_DPB1_macrophage
TCCCATPTAAGCGGTAGCTACCTTGCGGATCTCCCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

TACAGCCGTGAGCCACT

> P3_consensus_DPA1_capillary
TCCCATPTAAGCGGTAGCTACCTTGCGGATCTCCCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

HLA-B (Figures 3D and S4D)
q value: 2.7e-05
binomial p value: 1.7e-25.

> anchor
TGAGGACGCGAACAACAGATCTTCAAA

> target_1_HLA-B
AGAGCCTGCGGAAACCTGCGCCGC

> target_2_HLA-B
AGAGCCTGCGGAAACCTGCGCCGC

> consensus_1_HLA-B
TGAGGACGCGAACAACAGATCTTCAAA

> consensus_2_HLA-B
TGAGGACGCGAACAACAGATCTTCAAA

human Ig-kappa C-region (Figure 4B)
q value = 1.6E-35

> anchor
TGCGGGGAAAGATGGAAGACAGATCTTGC

> Targ0
GCTTGGTCCCTGCGCAAAAGTCCCGG

> Targ1
GCTTGGTCCCTGCGCAAAAGTCCCGG

> Targ2
GCTTGGTCCCTGCGCAAAAGTCCCGG

> Targ3
GCTTGGTCCCTGCGCAAAAGTCCCGG

CAGTTGCTGAAACAGCATACCTCTGACAGAAGCTTTGGAAGAATAAGAGCCAAGTGGGAAAGCTTTTCATCTTTGGCTTGCATTTCTG

OPEN ACCESS
>Targ4
GCTTGGTCCCTGGCCAAAAGTGTCGT
>Targ5
GCTTGGTCCCTGGCCAAAAGTGCCCG
>Targ6
CTTTGGTCCAGGCGAAAAGTGAATA
>Targ7
CTTTGGTCCCTGGCCGAAGCTCCACC

**human TCR-alpha C-region (Figure 4B)**

q value = 3.4E-5

>anchor
GTACACGGCAGGGTCAGGGTTCTGGAT
>Targ1
TGCCCTTGCCGAAGTTGAGTGCATACC
>Targ2
TCCCTGTACCAAGATTATCTTGGAAG
>Targ3
TGCCCTGTACCAAGATTATCTTGGAAG
>Targ4
TCCCTGTACCAAGATTATCTTGGAAG
>Targ5
TCCCCTCTTCCAAGAGCGCTTTCTGC
>Targ6
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ7
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ8
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ9
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ10
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ11
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ12
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ13
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ14
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ15
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ16
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ17
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ18
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ19
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ20
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ21
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ22
TTCCCTCTTCCAAGAGCGCTTTCTGC
>Targ23
TTCCCTCTTCCAAGAGCGCTTTCTGC
lemur Ig-heavy V-region (Figure 4B)
q value = 1.3E-11

>anchor
AGCCTGGGGGGTCCCTGAGACTCTCCT
>Targ0
AGTGACTACTACATGAGCTGGGTCCGC
>Targ1
AGCGACTATGGGATGAACTGGGTCCGC
>Targ2
AGCAACTCTTGATGACCTGGGTCCGC
>Targ3
AAGAACTATGAGATAAACTGGGTCCGC
>Targ4
AGCAGCTACTACATGCACTGGGTCCGC
>Targ5
AGTGACTACTACATGAACTGGGTCCGC
>Targ6
AGTCGACTACATGAACTGGGTCCGC
>Targ7
AGCAGCCATGGAATGCACCTGGGTCCGC
>Targ8
AGCAGCTACGATATGACCTGGGTCCGC
>Targ9
AGCAGCTATGACCTGGGTCCGC
>Targ10
AGTGACCACCACTAGCTGGGTCCGC
>Targ11
AGTGACTACCTTCATGGAACCTGGGTCCGC
>Targ12
AGCAGCTATGCCATGGAACCTGGGTCCGC
>Targ13
AGTAGTTATGGCATGAACTGGGTCCGC
>Targ14
AGTTACTATGGCATGAACTGGGTCCGC
>Targ15
ACCAATTGATGGAACCTGGGTCCGC
>Targ16
AGCAGCTATGGGATGCACTGGGTCCGC
>Targ17
AGCAGCTATGGGATGCACTGGGTCCGC

lemur TCR-alpha C-region (Figure 4B)
q value = 4.1E-7

>anchor
TCAGCTGGATACAGGCCGTCAGGGTG
>Targ0
AGTCTGGTCCTGGCTCAGGACCA
>Targ1
AGCCTGGTCCTGGCTCAGGACCA
>Targ2
AGCAGGTGCTCCGATTCACAAAGATGAGC
>Targ3
ACGGTGCTCCGATTCACAAAGATGAGC
>Targ4
AGTTGGGTGCCAGTTCCAAACACGGGT
>Targ5
AACTGGGTCCCGGATCCAAAGGTCAGT
>Targ6
AGGAGCTTCCTTTTCCCCAAGATGAGC
>Targ7
AGTGACTACCCATGGAACCTGGGTCCGC
>Targ8
AGCAGCTATGCCATGGAACCTGGGTCCGC
>Targ9
AGCAGCTATGCCATGGAACCTGGGTCCGC
>Targ10
AGTGACCACCACTAGCTGGGTCCGC
>Targ11
AGTGACTACCTTCATGGAACCTGGGTCCGC
>Targ12
AGCAGCTATGCCATGGAACCTGGGTCCGC
>Targ13
AGTAGTTATGGCATGAACTGGGTCCGC
>Targ14
AGTTACTATGGCATGAACTGGGTCCGC
>Targ15
ACCAATTGATGGAACCTGGGTCCGC
>Targ16
AGCAGCTATGGGATGCACTGGGTCCGC
>Targ17
AGCAGCTATGGGATGCACTGGGTCCGC
Octopus bimaculoides Myo-VIa (Figure 5A)
q value = 4.0e-03
(reverse-complements shown in Figure 5A)
>anchor
CCATTTTTGCTTTTTGTTTAAAATCCA
>target_1
ATTATATCACAAGTTATAAGGCATGCC
>target_2
ATTATATCTTAATAAATGGGACACTA

fucoxanthin chlorophyll a/c protein, diatom (Figure 5C)
q value = 6.0e-08.
(reverse-complements shown in Figure 5C)
>anchor
AAGTATCACAACGGAAGCGATGGAG
>target_1 (abundance order = 1)
ATACGTCCGCTCGTTGACCTCAGAAAT
>target_2 (abundance order = 6)
ATACGGCCGTCTCGTGACCTCAGAAAT
>target_3 (abundance order = 2)
ATACGTCCGCTCGTTGACCTCAGAAAT
>target_4 (abundance order = 4)
ATACGTCCGCTCGTTGACCTCAGAAAT
>target_5 (abundance order = 5)
ATACGTCCGCTCGTTGACCTCAGAAAT
>target_6 (abundance order = 3)
ACACGTCCATGCTTAATTTCGACATAT

Zostera marina NADPH quinone oxidoreductase subunit L (NdHL) (Figure 5D)
q value = 6.5e-56.
(reverse-complements shown in Figure 5D)
>anchor
AATCGAAGCCAATTCATGATGATAGGC
>target1
GGCATGATAAGGAAGTAGAAGAAAGCA
>target2
GGCATGATAAGGAAGTAGAAGAAAACA
>target3
GGCATGACAAGGAAGTAGAAGAAAGCA
>target4
TTCGATCATGCAGTTCAATCAATGATC

human MYL6 (Figure S4A)
>P2_anchor
AAGGTCCTCAGCCATCGACCCCATGTCGAC
>P2_consensus1_macrophage
GGCAAGCTGTCTTCTATAGTGAAGCCACATGTGCTTCCTCTGCTAGGATTTTGATGACATGATTGCCCGACTCTTCTCTCCTR
>P3_anchor (same as P2)
AAGGTCCTCAGCCATTCAGCACCATGC

>P3_consensus_macrophage
AAGGTCCTCAGCCATTCAGCACCATGCGGACGAGCTCTTCATAGTTGATACAACCATTGCTGTCCTCATGCCCTGCCACCAGCATCTCTACTTCTTC

>TTCCTCTGTCATCTTCTCACCCAGTGTGACAAGAACATGCCGGA

>P3_consensus_capillary
AAGGTCCTCAGCCATTCAGCACCATGCGGACGAGCTCCGCCCCATGGGCCCGTCACCCCGACAGGATATGCCTCACAAACGCTTCATAGTTGAT

CAACCATTGCCCTGTCCTCGACCAGCATCTCTACTTCTTCTCTT

>P3_target1
CAACCATTGCCCTGTCCTCATGCCCTGCC

>P3_target2
CGTCACCCCCGACAGGATATGCCTCACAA

mouse lemur COX2 (cytochrome c oxidase subunit II) (Figure S5A)
(reverse-complements are shown in Figure S5A)

>anchor
ATTTAGGCGCCCTGGGATAGCATCTGT

>target_1
TTTATGAATGTAGTACGTCTTCTGAAG

>target_2
TTTATGAATGTAGTACGTCTTCTGAAG

lemur IGLC3 with 97 targets (Figure S5B)

>anchor
ACCGAGGGGCGGCCTGGGCTGACCT

>Targ0
GCCGAACACCCAGTGCCACCACCTCCT

>Targ1
GCCGAAGATATGACCACCTCGGTGTC

>Targ2
GCCGAACACATGATTGTAGCTGCCATC

>Targ3
GCCGAATACATTAACACCACCTGTTGTC

>Targ4
GCCGAACACATAAACCACCTGTTGTC

>Targ5
GCCGAACACACCACCTGCTGCCACC

>Targ6
GCCGAACACATTAACACCACCTGTTGTC

>Targ7
GCCGAATACAGCAGCCTGGGCTGACCT

>Targ8
GCCGAAGATATGAGTTCTTCGCCACAC

>Targ9
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ10
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ11
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ12
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ13
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ14
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ15
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ16
GCCGAACACACCCAGACCCACCTGCTGTC

>Targ17
GCCGAACACACCCAGACCCACCTGCTGTC
GCCGAATACATAAGCACTCAGGCTTTT
>Targ18
GCCGAACCCGCCACTCAGGCTGCT
>Targ19
GCCGAATACATAAGCACCCTAGGCTGCT
>Targ20
GCCGAAGATATACCGCAGTACGCTACT
>Targ21
GCCGAACCTGACCACACTCAGGCTACT
>Targ22
GCCGAACCAACACCACCCACCTG
>Targ23
GCCGAACCACTGACGCAGCCTACT
>Targ24
GCCGAACCCAGCGACTGAGCTGC
>Targ25
GCCGAACCTGACCACACTCAGGCTACT
>Targ26
GCCGAACACATCGACCTGAGCTGC
>Targ27
GCCGAACACATGACCCACCTG
>Targ28
GCCGAAGATATTAGCACACTG
>Targ29
GCCGAAGATATACCTACCTGAGCTGC
>Targ30
GCCGAACCTAACTCAGCTAGGCC
>Targ31
GCCGAACACATACGCTGCTTG
>Targ32
GCCGAACACATGACCCACCTG
>Targ33
GCCGAACCCAGCGACTGAGCTGC
>Targ34
GCCGAACCCAGCGACTGAGCTGC
>Targ35
GCCGAACCTGACCACACTCAGGCTACT
>Targ36
GCCGAACACATCGACCTGAGCTGC
>Targ37
GCCGAACACATCGACCTGAGCTGC
>Targ38
GCCGAACACATGACCCACCTG
>Targ39
GCCGAACACATCGACCTGAGCTGC
>Targ40
GCCGAACCCAGCGACTGAGCTGC
>Targ41
GCCGAACACATGACCCACCTG
>Targ42
GCCGAACCCAGCGACTGAGCTGC
>Targ43
GCCGAACACATGACCCACCTG
>Targ44
GCCGAACACATGACCCACCTG
>Targ45
GCCGAACACATGACCCACCTG
>Targ46
lemur TCR-beta J-region (Figure S5C)

>anchor
CGGGGCTCCCTGGGCCCCAGAAGACTGTCTC
>Targ0
TGCCGCTGCGAGTGTAGACGCCGCTGT
>Targ1
CGCAGAGATACAGGGCCGAGTCCCCCA
>Targ2
TGGCACAGAGGTACGTGGCGGAGTCTT
>Targ3
TGCTGGCACAGAGGTACGTGGCAGAGT
>Targ4
AGAGGAACAGGGCCGAGTCCCCCAGCG
>Targ90
GCCGAACACACACTGCTGCCATC
>Targ91
GCCGAATACAAATGGTTGCTGAACCAC
>Targ83
GCCGAACACATGACTGTAGCTGTCATCATA
>Targ82
GCCGAACACACTGTAGCTGTCATCATA
>Targ84
GCCGAACACACTGCTGCC
>Targ80
GCCGAACACACACTGCTGCC
>Targ86
GCCGAACACACTGCTGCC
>Targ81
GCCGAACACACTGCTGCC
>Targ87
GCCGAACACACTGCTGCC
>Targ88
GCCGAACACACTGCTGCC
>Targ89
GCCGAACACACTGCTGCC
>Targ90
GCCGAACACACTGCTGCC
>Targ91
GCCGAACACACTGCTGCC
>Targ92
CAGAGGCCTGTGCCACCTGGGGAGCC
>Targ93
GCCGAACACACTGCTGCC
>Targ94
GCCGAACACACTGCTGCC
>Targ95
GCCGAACACACTGCTGCC
>Targ96
GCCGAACACACTGCTGCC

lemur TCR-gamma V-region (Figure S5D)

>anchor
CCGGTCCCTGGGCCCCAGAAGACTGTCTC
>Targ0
TGCCGCTGCGAGTGTAGACGCCGCTGT
>Targ1
CGCAGAGATACAGGGCCGAGTCCCCCA
>Targ2
TGGCACAGAGGTACGTGGCGGAGTCTT
>Targ3
TGCTGGCACAGAGGTACGTGGCAGAGT
>Targ4
AGAGGAACAGGGCCGAGTCCCCCAGCG
ACCCTCACCATTCAAAATGTAGAGAAA
>Targ0
TGCCCTGGAACCTTTCAATGAATGAAAC
>Targ1
TGCCCTCCTGGGACTGACGACTTACCAA
>Targ2
TGCCCTCCTGGGAAGTTGAATTTTTATAG
>Targ3
TGCCCTCCTGGGAGTTGCACAGTGTCAC
>Targ4
GCCCGTGAACTCTTCAGTAATGGAAC
>Targ5
TGCCCTCCTGGGAGTCTAGGAAACTCTT
>Targ6
TGCCCTCCTGGGAGTTGCACAGAAGATT
>Targ7
TGCCCTCCTGGGAGTTGCACAGAAGATT

**Octopus bimaculoides carboxypeptidase D** (Figure S6A) (reverse-complements are shown in Figure S6A)
>anchor
GGAATTAGAAGAAAAATCTATTATGAA
>target_1
AAATGTTTAGGCCAATATCTAAAGGCA
>target_2
AAATGTTTAGGCCAATATCTAAAGGCA

**Octopus bimaculoides Upf2** (regulator of nonsense transcripts 2) (Figure S6B) (reverse-complements are shown in Figure S6B)
>anchor
GTATTGCACTGCATTGTACTGCACTGT
>target_1
CGCTGCTGCTGCTGCTGCTGCCAATTG
>target_2
CGCTGCTGCTGCTGCTGCCAATTGCCT

**Octopus bimaculoides netrin receptor/DCC** (Figure S6C) (reverse-complements are shown in Figure S6C)
>anchor
TCTATTACAGCTATCATCAATACACTT
>target_1
TTGGATGTCTTCGTGTTCTCACTGCAG
>target_2
TTGGATGTCTTTGTGTTCTCACTGCAG

**HMG-box** (diatom) (Figure S7A) (reverse-complements are shown in Figure S7A)
>anchor
TGCGTCCCTGGAATTTCTGCTTCTTCTT
>target_1
TATCCGAAAAGGCCCTCCCACATTTCAC
>target_2
CGTCCGTAGACAGTCTCCCCACATTTCAC

**ferredoxin** (diatom) (Figure S7B) (reverse-complements are shown in Figure S7B)
>anchor
ACGGCACAGTAGGGAATTTCTGCAATTCC
>target_1
GGCTTCTTCAGAGTCGACAGAATGAA
>target_2
GGCTTCTTCAGAGTCGACAGAATGAA
QUANTIFICATION AND STATISTICAL ANALYSIS

SPLASH p values
SPLASH’s analysis is based on a new statistical method for analyzing contingency tables, to reject the null hypothesis that row (target) counts are drawn from the same distribution for all samples. Despite its rich history, the field of statistical inference for contingency tables still has many open problems.80 The field’s primary focus has been on either small contingency tables (2 \times 2, e.g., Fisher’s exact test81), high counts settings where a chi-square test yields asymptotically valid p values, or computationally intensive Markov Chain Monte Carlo (MCMC) methods. While contingency tables have been widely analyzed in the statistics community,80,82,83 to our knowledge no existing tests provide closed form, finite-sample valid statistical inference with desired power for the application at hand. In this work, we develop a flexible statistical test that provides closed form p value bounds, meaning that no permutation, resampling or numerical solutions to complex likelihoods are required for significance tests. In subsequent work, more sophisticated (optimization-based) approaches to computing improved \( f \) and \( c \) have been developed, leveraging a linear algebraic perspective on the test statistic.52

We develop a test statistic \( S \) that has power to detect many forms of sample-dependent sequence variation and is designed to have low power when there are a few outlying samples with low counts. Considering the target \( \times \) sample contingency table (\( I \times J \)), we first randomly draw a vector \( f \), which maps each of the \( I \) targets independently to \( \{0,1\} \) equiprobably. We then compute the mean value of targets with respect to this function. Next, we compute the mean target value within each sample with respect to \( f \).

Then, an anchor-sample score is constructed for sample \( j \), \( S_j \), as a scaled version of the difference between these two (scaled by the square root of the total counts for this sample). Finally, the test statistic \( S \) is computed as the weighted sum of these \( S_j \), with weights \( c_j \) (which denote class-identity in the two-group case with metadata and are chosen randomly without metadata, see below).

In the below equations, \( Z_{j,k} \) denotes the target index (1 through \( I \)) of the \( k \)-th target observed for the \( j \)-th sample (for analysis purposes, we assume that the targets are drawn in a random order), \( n_j \) is the number of observations in this sample, and \( M \) denotes the total number of observations of this anchor.

\[
\hat{\mu} = \frac{1}{M} \sum_{j,k} f_{j,k}
\]

\[
\hat{\mu}_j = \frac{1}{n_j} \sum_{k=1}^{n_j} f_{j,k}
\]

\[
S_j = \sqrt{n_j} (\hat{\mu}_j - \hat{\mu})
\]

\[
S = \sum_{j=1}^{J} c_j S_j
\]
Statistically valid p value bounds (non-asymptotic, providing control of Type I error for finite number of observations) are computed as:

\[
P = 2 \exp \left( -\frac{2(1-a)^2\epsilon^2}{\sum_j c_j^2} \right) + 2 \exp \left( -\frac{2a^2Ms^2}{\left( \sum_j c_j \sqrt{\nu_j} \right)^2} \right) \quad \text{where} \quad a = \left( 1 + \frac{M \sum_j c_j^2}{\left( \sum_j c_j \sqrt{\nu_j} \right)^2} \right)^{-1}
\]

by applying Hoeffding’s inequality on these sums of independent random variables (under the null). We provide a graphical description of this procedure in Figure S1A, and details the derivation below.

First, observe that \( S \) has mean 0 under the null hypothesis. This allows us to bound the probability that the random variable \( S \) is larger than our observed anchor statistic as follows. Since \( f \) and \( c \) are fixed (independent of the data), and \( f(Z_{j,k}) \) are bounded between 0 and 1, we can apply Hoeffding’s inequality for bounded random variables. Defining \( \mu \) as the expectation with respect to the common underlying distribution of \( Z_{j,k} \) (unknown), we center our random variables by subtracting the sample mean \( \bar{\mu}_m \), our estimate of the true mean \( \mu \). Standard bounds can now be applied to decompose this deviation probability into two intuitive and standard terms, which we then bound with Hoeffding’s inequality for bounded random variables:

1) the probability that the statistic \( \tilde{S} \), constructed with unavailable knowledge of the true \( \mu \), is large

\[
\tilde{S} = \sum_j c_j (\mu_j - \mu)
\]

2) the probability that \( \bar{\mu} \) is far from \( \mu \).

\[
P(|S| \geq \epsilon) \leq P \left( \sum_{j, k} c_j f(Z_{j,k}) - \mu \geq \epsilon \right)
\]

\[
= P \left( \sum_{j, k} c_j f(Z_{j,k}) - \mu \geq (1-a)\epsilon \right) + P \left( \sum_{j} c_j \sqrt{\nu_j} \geq a\epsilon \right)
\]

\[
\leq \min_{a \in (0, 1)} \left[ P \left( \sum_{j, k} c_j f(Z_{j,k}) - \mu \geq (1-a)\epsilon \right) + P \left( \sum_{j} c_j \sqrt{\nu_j} \geq a\epsilon \right) \right]
\]

\[
= \min_{a \in (0, 1)} \left[ 2 \exp \left( -\frac{2(1-a)^2\epsilon^2}{\sum_{j, k} c_j^2} \right) + 2 \exp \left( -\frac{2a^2Ms^2}{\left( \sum_j c_j \sqrt{\nu_j} \right)^2} \right) \right]
\]

The bound is optimized to within a factor of 2 by equating the two terms, with \( a \) as stated.

This statistic is computed for \( K \) different random choices of \( f \), and in the case where sample group metadata is not used or available, jointly for each of the \( L \) random choices of \( c \) (each entry drawn independently as \( \pm 1 \)), here with \( K = 10 \) and \( L = 50 \). The choice of \( f \) and \( c \) that minimize the p value bound are reported, and are used for computing the p value bound for this anchor. To control Type I error we apply Bonferroni correction over the \( L \times K \) multiple hypotheses tested (just \( K \) when sample metadata is used and randomization on \( c \) is not performed). Then, to determine the significant anchors, we apply Benjamini-Yekutieli (BY) correction\(^84\) to the list of p value bounds (one for each anchor), yielding valid FDR controlled q values reported throughout the manuscript implemented with the statsmodels.api.stats.multipletests functionality in python. Additional theoretical properties of this statistic, and an improved p value bound, have been developed in follow-up work.\(^52\)
**SPLASH effect size**

SPLASH provides a measure of effect size when the $c_j$’s used are $\pm 1$, to allow for prioritization of anchors with large inter-sample differences in target distributions (as the $p$ value bound is heavily impacted by the number of observations). Effect size is calculated based on the split $c$ and vector $f$ that yield the smallest SPLASH $p$ value bound. Fixing these, the effect size is the absolute value of the difference between the mean target value (with respect to $f$) across those samples with $c_j = +1$ denoted $A_+$, and the mean target value (with respect to $f$) across those samples with $c_j = -1$ denoted $A_-$. 

\[
\left| \sum_{j \in A_+} \frac{1}{n_j} \bar{y}_j - \sum_{j \in A_-} \frac{1}{n_j} \bar{y}_j \right|
\]

The effect size should be thought of under the alternative hypothesis where the columns follow multinomial distributions with probability vector $p_1$ or probability vector $p_2$, depending on the group identity $c_j$. The effect size we compute can be thought of in this scenario as measuring the difference between the expectation of $f$ under $p_1$ and $p_2$. In the case of maximizing the effect size over all possible $0,1$-valued $f$, the effect size will be equal to the total variation distance between the empirical distributions of the group $c_j = +1$ and $c_j = -1$. Thus, the effect size will be 1 if and only if the two sample groups partition targets into 2 disjoint sets on which the function $f$ takes opposite values, as to be expected from the total variation distance interpretation (Figure S1B). This $f$ will place a value of 1 on targets where the empirical frequency of the +1 group $p_{1,t}$ is larger than that of the −1 group $p_{2,t}$. Since $p_1$ and $p_2$ are probability distributions, this ends up being exactly the total variation distance between them (i.e., half the vector $k_1$ distance).

**Binomial p value bound computation for plots depicting target fraction abundance**

We provide $p$ value bounds to quantify the visually striking nature of the plots depicting fraction abundance of a specific target (target 1 by default). Under a null model, where all samples are expressing this target with the same probability, the number of times each sample expresses target 1 is binomial($n_j$, $p$), for common $p$. As seen from the plots, many samples exhibit highly deviating occurrences (number of observations of target 1 that are far from the expected $p n_j$). The $p$ values we provide to this effect are not used in any SPLASH discovery or analysis, and are just used to quantify the visually striking nature of these plots.

$p$ value bounds are constructed as follows: first, we compute $p$, the average occurrence of target 1 for this anchor (sum of counts of observations of target 1 divided by the total number of observations). Then, for all possible $n_j$, we compute 1% and 99% quantiles (confidence bounds) for a binomial distribution with $n_j$ trials and heads probability $p$. If the fraction of target 1 in each sample was independent of sample identity, and were indeed binomially distributed, then each sample would have at least a 98% probability of falling within this confidence interval. Thus, we compute our test statistic $X$ as the number of samples that fall outside of the [1,99] quantiles, and compute as our $p$ value bound the probability that a binomial random variable $\text{Bin}(m,q) \geq X$, where with $m =$ number of deviating samples and $q = 0.02$.

While intuitive, the above analysis is loose. Firstly, since binomials are discrete distributions, we will rarely be able to compute exact 1% and 99% quantiles. Thus, the probability that for any given $n_j$ a sample will fall outside of the $[1,99]$ quantiles, which we denote $q_j$, is almost always substantially less than 0.02. The true distribution of $X$ is then poisson binomial, with this vector of probabilities (all at most 0.02), one for each sample. However, as this $p$ value is numerically difficult to compute, we bound this $p$ value as the probability that $\text{Bin}(m,q') \geq X$, where $m =$ number of samples with $q' = \max_j q_j$, where $q' \leq 0.02$. 

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**A  p-value computation**

| \( t \) | 1 | 0 | 1 | 0 |
|---|---|---|---|---|
| 45 | 5 | 2 | 1 |
| 93 | 11 | 3 | 3 |
| 50 | 6 | 1 | 2 |
| 2 | 38 | 1 | 1 |
| 20 | 370 | 7 | 8 |

| \( \hat{\mu}_j \) | .89 | .87 | .86 | .07 | .07 |
|---|---|---|---|---|---|
| \( S_j \) | 4.0 | 5.6 | 4.0 | -1.7 | -5.4 |
| \( c \) | 1 | 1 | 1 | 1 | 1 |

1) Compute \( \hat{\mu}_j \)  
2) Compute \( S_j = \sqrt{n_j}(\hat{\mu}_j - \bar{\mu}) \)  
3) Compute \( S = \sum c_j S_j \)  
4) Concentration inequality  
5) BY correction across anchors for MHT

**B  effect size computation**

| \( c' \) | 1 | 0 | 1 | 0 |
|---|---|---|---|---|
| 45 | 5 | 2 | 1 |
| 93 | 11 | 3 | 3 |
| 50 | 6 | 1 | 2 |
| 2 | 38 | 1 | 1 |
| 20 | 370 | 7 | 8 |

1) Compute \( \hat{\mu}_1 = 198, 22, 6, 6 \)  
2) Compute \( \hat{\mu}_2 = 22, 408, 8, 9 \)  
3) Compute \( f^* \) value

\[ \hat{\mu}_1 = \frac{204}{232} = 0.88 \]  
\[ \hat{\mu}_2 = \frac{30}{417} = 0.067 \]

Effect Size = \( \hat{\mu}_1 - \hat{\mu}_2 = 0.81 \)

**Figure S1.** SPLASH computations, related to Figure 1

(A) p value bound computation for SPLASH for fixed vectors \( t \) and \( c \). Contingency table transposed for visual convenience (rows are samples, and columns are targets). Starting with a counts matrix, SPLASH utilizes the vector \( t \) to map targets to values within \([0,1]\). The mean with respect to \( t \) is taken over the targets in each sample \( j \) to yield \( \hat{\mu}_j \), and an estimate for the mean over all target observations of \( t \) is taken, yielding \( \bar{\mu} \). The anchor-sample scores \( S_j \) are then constructed as the difference between the row mean \( \hat{\mu}_j \) and the overall mean \( \bar{\mu} \), scaled by \( \sqrt{n_j} \) (the number of observations in sample \( j \)). These anchor-sample scores are weighted by \( c_j \) and summed to yield the anchor statistic \( S \). Finally, a p value bound is computed utilizing classical concentration inequalities, which we correct for multiple hypothesis testing (with dependence) by constructing \( q \) values using Benjamini-Yekutieli, a variant of Benjamini-Hochberg testing that corrects for arbitrary dependence.

(B) Effect size computation for SPLASH. Reported effect size is calculated based on the split \( c \) and vector \( t \) that yielded the most significant SPLASH p value. Fixing these, the effect size is computed as the difference between the mean across targets (with respect to \( t \)) across those samples with \( c_j = +1 \) and the mean across targets (with respect to \( t \)) across those samples with \( c_j = -1 \). This should be thought of as studying an alternative where samples from \( c_j = +1 \) have targets that are independent and identically distributed with mean (under \( t \)) \( \mu_1 \) and samples with \( c_j = -1 \) have targets that are independent and identically distributed with mean (under \( t \)) \( \mu_2 \). The total effect size is estimated as \( \hat{\mu}_1 - \hat{\mu}_2 \), which relates to the total variation distance between the two target distributions (STAR Methods).
We performed SPLASH protein domain profiling in a dataset of virally enriched samples from breakthrough infections in patients that had been vaccinated against rotavirus; nearly all had rotavirus infection, but some also had other coinfections. The top domains were rotavirus VP3 (Rotavirus_VP3, 76 SPLASH hits vs. 9 control hits) followed by rotavirus NSP3 (Rota_NSP3, 87 SPLASH vs. 35 control hits), indicating that these rotavirus genes have sequences that vary significantly among these patients; they have roles in blocking host innate immunity, and so may be under selective pressure for variation.
Figure S3. Unsupervised SPLASH recovers supervised results, SPLASH data size reduction, related to Figures 1 and 3

(A) SPLASH’s unsupervised approach (selection of random $c$ vectors) yields good performance and is able to recapitulate results obtained by the supervised method. We validate this on the HLCA dataset, where given metadata indicate whether a sample is from macrophage or capillary cells, which the supervised approach utilizes. Of the 3,336 anchors (848 genes) called by the supervised approach in individual P2 (BY correction, alpha = 0.05), 71% of the genes called were also called by SPLASH’s unsupervised approach (6,407 called anchors, 1,503 genes). The left plot indicates how many times each gene (dot) was called by unsupervised vs. supervised approaches. Red dots indicate those genes not called by SPLASH’s unsupervised approach. The right plot shows the fraction of genes that are called at least $x$ times by the supervised approach that are also called by the unsupervised one. For $x = 2$ (i.e., all genes hit by >2 anchors), our unsupervised approach calls >92% of those genes called by the supervised one.

(B) If reference alignment is desired, aligning only SPLASH results (consensuses) reduces the amount of data for alignment by >500-fold compared with aligning all the raw sequencing reads; this is consistent in two different datasets. MiB = mebibyte = 1,048,576 bytes.
Figure S4. Cell-type-specific expression of MYL6, related to Figure 3
(A) MYL6 SPLASH results from two individuals (P2 and P3) are shown. For all the plots, columns represent different samples (single cells). The heatmap with rows labeled by target sequences is the sample x target contingency table; entries indicate the respective target fraction for a given sample, x ticks give the anchor counts in that sample, and y ticks give the target counts across all samples. The top two plots show \( c_j \)’s; “sample sheet \( c_j \)” are the cell-type labels provided in the original study (pink, macrophage; blue, capillary), and “\( c_j \) found by random” are SPLASH’s unsupervised assignments. The two bottom plots relate to the consensus sequences. The lower left plot depicts the nucleotide sequence using colors for the bases. Each column corresponds to the consensus sequence for the sample of the same column in the contingency table. The rows are base pair positions along each consensus. These consensus sequences are variable length, and a value of \( \frac{1}{C_0} \) (yellow color) on the bottom of a sequence indicates that the consensus has ended. The bottom right plot shows the fraction agreement per nucleotide within a sample with its consensus sequence. For samples where only one isoform/SNP is expressed, the consensus stays near 100%, while for samples with a diverse set of targets, the consensus is less uniform.
(B) Macrophages tend to express the exon-skipping isoform, as shown by dot plots, with each dot representing the relative isoform expression in a single cell; a bar marks the average relative expression.
(C) UCSC Genome Browser diagram showing BLAT mapping of macrophage and capillary consensus sequences, which fully cover inclusion or exclusion of the short alternative exon.
Figure S5. Lemur COX2 allelic detection, additional Ig/TCR anchors, related to Figure 4

For (B)–(D), target × sample heatmaps are shown on the left (samples are individual cells), and bp color maps are on the right. The small numbers at the sides of the heatmaps are summed target counts (over rows or columns).

(A) COX2. SPLASH detects numerous anchors that have target fractions that correspond precisely to the identities of the two lemur individuals represented in the set of cells analyzed. One example is an anchor in COX2. The alignment of two consensuses (both from NKT cells) shows that targets 1 and 2 differ at a single silent position (C vs. T) and another silent SNP is also present outside the target region (G vs. A); these positions are highlighted in red. Consensuses 1 and 2 align perfectly to different lemur mitochondrial genome accessions (NC_028718.1 and KR911907.1). Targets 1 and 2 are found exclusively in individuals L4 and L2, respectively.

(B) Ig-lambda C region. Depicted is a lemur anchor that maps to the 5' end of the human Ig-lambda C3 segment. It has 97 different targets, which lie within the hypervariable CDR3 region. Nearly all targets are expressed clonotypically (i.e., cells do not share targets).

(C) TCR-beta J region. We analyzed NKT cells from lemur. In humans and mice, a subset of NKT cells express stereotypical or shared TCR genes; Figure 4B shows an example anchor in TCR-alpha. Here, we show an anchor that maps to a human TCR-beta J region (J2-1); targets reside in the V region. A number of the NKT cells express a shared TCR-beta target (seen in the top row of the heatmap). All of these cells derive from individual L4.

(D) TCR-gamma V region. There is evidence for shared TCR-gamma sequences in NKT cells as well. This anchor maps to a human TCR-gamma V region (TRGV8); the targets lie at the V-J junction. Individual L4 has two different shared targets (rows 1 and 3), while individual L2 shows three cells expressing a shared target (row 2). Other NKT cells express unique targets, however.