A cellular taxonomy of the adult human spinal cord

Graphical abstract

Highlights
- Single-nucleus RNA sequencing and spatial transcriptomics of adult human spinal cord
- Glial classes include proliferative microglia and white and gray matter astrocytes
- Dorsal neuron types are more discrete and ventral neuron groups are more overlapping
- The human motoneuron transcriptional profile is enriched for genes related to ALS

Authors
Archana Yadav, Kaya J.E. Matson, Li Li, ..., Claire E. Le Pichon, Vilas Menon, Ariel J. Levine

Correspondence
vm2545@cumc.columbia.edu (V.M.), ariel.levine@nih.gov (A.J.L.)

In brief
Yadav, Matson, et al. use single-nucleus RNA sequencing, spatial transcriptomics, and immunohistochemistry to profile the cell types of the adult human spinal cord, identifying 64 glial and neuronal populations. This resource reveals how the unique molecular environments of specific cell types could contribute to chronic pain or neurodegeneration.
A cellular taxonomy of the adult human spinal cord

Archana Yadav,1,16 Kaya J.E. Matson,2,3,16 Li Li,2 Isabelle Hua,2 Joana Petrescu,1,4 Kristy Kang,1,4 Mor R. Alkaslasi,5,6 Dylan I. Lee,1 Saadia Hasan,7 Ahmad Galuta,2,10 Annemarie Dedek,8,9 Sara Ameri,9 Jessica Pameli,8,9 Mohammad M. Alshardan,8 Feras Abbas Qumqumji,9 Saud M. Alhamad,8 Alick Pingbei Wang,6 Gaetan Poulen,10 Nicolas Lonjon,1,10 Florence Vachiery-Lahaye,11 Pallavi Gaur,1 Mike A. Nalls,1,12,13 Yue A. Qi,12 Dragan Maric,14 Michael E. Ward,1 Michael E. Hildebrand,7,4 Pierre-Francois Mery,1,15 Emmanuel Bourinet,1,15 Luc Bauchet,1,15 Eve C. Tsai,6 Hemali Phatnani,1,4 Claire E. Le Pichon,8 Vilas Menon,1,17,1* and Ariel J. Levine2,17,1*,
1Department of Neurology, Center for Translational and Computational Neuroimmunology, Columbia University, New York, NY, USA
2Spinal Circuits and Plasticity Unit, National Institute of Neurological Disorders and Stroke, Bethesda, MD, USA
3Johns Hopkins University Department of Biology, Baltimore, MD, 21218, USA
4Center for Genomics of Neurodegenerative Disease, New York Genome Center, New York, NY, USA
5Unit on the Development of Neurodegeneration, Eunice Kennedy Shriver National Institute of Child Health and Human Development, Bethesda, MD, USA
6Department of Neuroscience, Brown University, Providence, RI, USA
7Inherited Neurodegenerative Diseases Unit, National Institute of Neurological Disorders and Stroke, Bethesda, MD, USA
8Neuroscience Program, Ottawa Hospital Research Institute, Ottawa, ON, Canada
9Department of Neuroscience, Carleton University, Ottawa, ON, Canada
10Department of Neurosurgery, Gui de Chauliac Hospital, and Donation and Transplantation Coordination Unit, Montpellier University Medical Center, Montpellier, France
11Laboratory of Neurogenetics, National Institute on Aging, National Institutes of Health, Bethesda, MD, USA
12Center for Alzheimer’s and Related Dementias, National Institutes of Health, Bethesda, MD, USA
13Data Tecnica International LLC, Glen Echo, MD, USA
14Flow and Imaging Cytometry Core Facility, National Institute of Neurological Disorders and Stroke; Bethesda, MD, USA
15Institute of Functional Genomics, Montpellier University, CNRS, INSERM, Montpellier, France
16These authors contributed equally
17Senior author
18Lead contact
*Correspondence: vm2545@cumc.columbia.edu (V.M.), ariel.levine@nih.gov (A.J.L.)
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SUMMARY

The mammalian spinal cord functions as a community of cell types for sensory processing, autonomic control, and movement. While animal models have advanced our understanding of spinal cellular diversity, characterizing human biology directly is important to uncover specialized features of basic function and human pathology. Here, we present a cellular taxonomy of the adult human spinal cord using single-nucleus RNA sequencing with spatial transcriptomics and antibody validation. We identified 29 glial clusters and 35 neuronal clusters, organized principally by anatomical location. To demonstrate the relevance of this resource to human disease, we analyzed spinal motoneurons, which degenerate in amyotrophic lateral sclerosis (ALS) and other diseases. We found that compared with other spinal neurons, human motoneurons are defined by genes related to cell size, cytoskeletal structure, and ALS, suggesting a specialized molecular repertoire underlying their selective vulnerability. We include a web resource to facilitate further investigations into human spinal cord biology.

INTRODUCTION

The mammalian spinal cord relays, processes, and transforms sensory inputs and descending cues from the brain into sensory, motor, respiratory, and autonomic outputs. These critical processes rely on a diverse array of spinal cord cell types, each with their own molecular repertoires, functions, and vulnerabilities to injury and disease. Most prominently, spinal motoneurons specifically degenerate in spinal muscular atrophy and amyotrophic lateral sclerosis (ALS), though the molecular basis for this selective phenotype is not clear.2,3 Spinal cord cell types have been extensively studied in model organisms, including molecular profiling at the single-cell level, to identify candidate cellular mechanisms for human pathophysiology including in chronic pain, neurodegeneration, and spinal cord injury.4-12 However, technical obstacles and limited access to high-quality tissue specimens have prevented the full application of single-cell approaches to study human spinal cord biology directly.
Thus, prior work has only been done on limited cell types or in human fetal tissue.13–15

To characterize the cell types of the adult human lumbar spinal cord, we used recently optimized tissue extraction methods on spinal cords from organ donor subjects and performed single-nucleus RNA sequencing (snRNA-seq) of more than 50,000 nuclei. We identified 64 unique clusters, including 29 non-neuronal populations and 35 neuronal populations, and validated many of the expression patterns with spatial transcriptomics. We established a comprehensive taxonomy of the neuronal clusters, compared them with their mouse counterparts, and created a publicly available, browsable interface as a resource for the field (https://vmenon.shinyapps.io/humanspinalcord/). Finally, we performed a focused analysis on the transcriptional profile of spinal motoneurons, identifying a molecular signature that could underlie their selective vulnerability in neurodegenerative disease.

RESULTS

We obtained postmortem lumbar spinal cord tissue from 14 donor transplant cases (Table S1), using neuroprotective conditions (see STAR Methods). For snRNA-seq experiments, nuclei were isolated and profiled from seven donors (Figure 1A), resulting in a dataset of 55,289 nuclei after quality control (Figure S1). Initial clustering of all nuclei clearly distinguished the major known cell classes, including oligodendrocytes and their precursors and progenitors, meningeal cells, astrocytes, endothelial and pericyte cells, microglia, and neurons; the latter included glutamatergic neurons, GABAergic/glycinergic neurons, and motoneurons (Figure 1B). To determine whether the overall proportions of cell classes that we observed in the sequencing dataset reflected in vivo tissue composition, we analyzed the prevalence of oligodendrocytes, astrocytes, microglia, and neurons in adult human lumbar spinal cord tissue using antibody staining with classic cell class markers. We found similar proportions for neurons, astrocytes, microglia, and oligodendrocytes (Figures 1C and 1D, p = 0.67, p = 0.33, p = 0.06, p = 0.06; Figure S1) in tissue compared with the snRNA-seq dataset. Overall, the major cell classes in the sequencing dataset showed clear segregation of previously reported markers for these cell types, thus allowing for further investigation within each of these broad classes (Figures 1B and S1–S6), as described below.

Glial and support cell populations of the adult human lumbar spinal cord

We re-clustered non-neuronal cells and identified 29 subpopulations. We further characterized these groups by inspecting expression of known marker genes and examined their spatial distribution using spatial transcriptomics on tissue from five donors (Figure 2).

Among oligodendrocytes and related populations, we observed two groups of Schwann cells, a population of oligodendrocyte precursor cells and related progenitors, and six populations of oligodendrocytes that were distributed over the entire spinal cord tissue with a bias for the white matter, as expected (Figures 2A–2D, 2Q, S7, and S8).

Among support cells, we identified four populations of meningeal-related cells that included putative meningeal fibroblasts (Meninges-1, SLC4A4) and perivascular fibroblasts (Meninges-3 and Meninges-4, expressing ABCA8, DCN, and COL1A1) as well as a group of ependymal cells (Figures 2E–2H, 2Q, S7, and S8).

Among astrocytes (“Astro”), we identified three populations, including one localized to the white matter (WM Astro) that expressed the fibrous astrocyte marker CD44. The other two astrocyte populations were localized to the gray matter (GM Astro-1 and GM Astro-2) and were enriched for genes involved in neural metabolism and signaling, including the GABA transporter SLC6A11, the AMPA receptor regulator SHISA9, and the synaptic adhesion protein TENM2 (Figures 2I–2L, 2Q, S7, and S8).

Among vascular cells, we identified two endothelial cell populations, one of which represented putative venous/capillary cells (Endothelial-1, expressing IL1R1, ACKR1, ABCG2, and MFSD2A), while the other represented putative arterial cells (Endothelial-2, expressing SEMA3G, BMX, VEGFC, and PLCG2). We also identified a putative pericyte population expressing PDGFRB and NOTCH3 that likely included vascular smooth muscle cells as well (which are marked by SLIT3, ACTA2, and MYH11) and a cluster of lymphocytes (Figures 2E–2H, 2Q, S7, and S8).

We observed six populations of microglia (“Micro”) and one population of macrophages. A population of putative perivascular microglia were similar to a previously described microglial population in mice.16 This population showed enriched expression of inflammatory-related genes (CCL3, CCL4, IL1B, and ATF3) and were found just outside the main artery at the midline of the ventral horn of the spinal cord. Similarly, the cluster of macrophages (marked by MRC1, F13A1, LYVE1, and CD163) were found near, and within, the region of this main vessel (Figures 2M–P, 2Q, S7, and S8). We also observed a putative proliferative type of microglia characterized by expression of POLQ, TOP2A, and MKI67 (Figures 2M–P, 2Q, S7, and S8). In prior work on healthy adult mouse spinal cord, proliferative microglia were not observed.17 We therefore performed antibody staining on postmortem tissue from three organ donor subjects not included in the snRNA-seq dataset to confirm the existence of this population in intact tissue. Indeed, we found that 23% of IBA1+ microglia in tissue co-expressed the proliferative marker Ki67 (Figure S9, 25% ± 0.5% of cells were IBA1+, with 5.8% ± 0.38% of cells double positive for IBA1 and Ki67). Although prior studies on postmortem and surgically resected adult human brain tissue have also identified proliferative microglia,18 it is yet to be determined whether this reflects normal adult human biology, is associated with the advanced age of the tissue donors, or is due to peri-mortal changes in spinal cord. Proliferative microglia were enriched in white matter but also interestingly were found in the ventral horn near putative motoneurons (Figure S9).

Neuronal atlas of the adult human lumbar spinal cord

To characterize the neuronal populations of the adult human lumbar spinal cord, we subclustered the neuronal nuclei and identified 35 populations. These included a large population of spinal motoneurons (described below), a cluster defined by expression of immediate-early response genes (IEGs), and 33 main populations of spinal neurons (Figures 3A and S10). Each cluster contained nuclei from all seven donors, with
Figure 1. A single cell catalog of the human spinal cord reveals gene expression signatures of major cell classes

(A) Lumbar spinal cord tissue was obtained from seven subjects and processed for snRNA-seq.

(B) UMAP plot showing the major cell classes of the human spinal cord. Cells of the oligodendrocyte lineage are shown in pink/purple and include two populations of Schwann cells (Schwann-1 and Schwann-2), oligodendrocyte precursor cells (OPCs), progenitors (Oligo Progen), and six groups of oligodendrocytes (Oligo-1 through Oligo-6). Microglia are shown in green and include a putatively proliferating population (Prolif Micro), five groups of microglia (Micro-1 through Micro-4 and Perivascular Micro), as well as a population of macrophages. Astrocytes are shown in turquoise and include three populations (WM Astro, GM Astro-1, and GM Astro-2). Meninges are shown in blue and include four populations (Men-1 through Men-4). Vascular cells are shown in teal and include two groups of endothelial cells (Venous/Capillary Endo and Arterial Endo) and pericytes/smooth muscle cells (Peri/SMC). Ependymal cells are shown in cyan. Neurons are shown in orange and include seven broad classes based on their neurotransmitter status and putative location: motoneurons (MN), excitatory dorsal neurons (ExDorsal), inhibitory dorsal neurons (InhDorsal), excitatory mid neurons (ExM), excitatory ventral neurons (EV), inhibitory mid neurons (InhM), and inhibitory ventral neurons (InhV).

(C) Bar plot showing the proportion of each cluster in each donor (n = 7). Error bars are ±SEM.

(D) Multiplexed immunohistochemistry of the lumbar human spinal cord, stained for NeuN (yellow), IBA1 (green), SOX9 (turquoise), and OLIG2 (pink). Brightfield (BF) is shown in white. Mean percent of DAPI+ cells expressing NeuN, OLIG2, IBA1, and SOX9 are noted in the bottom right corner of each inset (n = 2). Scale bars indicate 500 μm.

See also Figures S1–S6.
the exception of Ex-Dorsal-1, Inh-Dorsal-4, and Inh-Dorsal-5 (Figures 3A and S11; Table S2).

The three main axes of gene expression variability among human spinal neurons were genes related to motoneuron identity, to spatial location, and to neurotransmitter status (Figure S12).

To assign putative locations for each population, we used Cell2location predictions based on spatial transcriptomics data, the spatial distribution of RNA expression for key marker genes, and comparison with data from macaque and mouse, sorting clusters into general categories of dorsal, mid, and ventral cell types (Figures 3B–3D). We next assigned putative neurotransmitter status to each population, identifying 19 glutamatergic populations (defined by the expression of SLC17A6) and 14 GABA/glycinergic populations (defined by expression of GAD1, GAD2, PAX2, and SLC6A5). A dendrogram of the overall cluster relationships confirmed that location and neurotransmitter status were the major organizational axes of spinal neurons, with dorsal excitatory clusters defining the first branch-point, and then dorsal inhibitory clusters and mid-ventral clusters (of both general neurotransmitter types) defining subsequent branchpoints (Figure 3C). We therefore adopted a nomenclature for human spinal neuron populations that references both neurotransmitter status and location. These and other cell type features are summarized in Table S3, with selected data in Figures 3 and S10–S13.

Dorsal excitatory neurons were organized into 12 distinct clusters mainly localized to the superficial dorsal horn (lamina I/II), with the exception of Ex-Dorsal-4, which was localized to lamina III (Figures S7 and S8). Expression of LMX1B, as well as LBX1 and TLX3, in many of these populations suggested a dl5/dl5B embryonic origin for these cells (Figure S13). These populations all expressed the vGLUT2 gene SLC17A6, with Ex-Dorsal-6 additionally expressing the vGLUT3 gene SLC17A8. One major group (Ex-Dorsal-1, Ex-Dorsal-2, and Ex-Dorsal-4) was likely non-peptidergic and distinguished by the MAF/MAFA bZIP transcription factors, while the other clusters were likely peptidergic, distinguished by genes such as PAM, TAC1, TAC3, NMU, and GRP (Figure S10). The dorsal inhibitory neurons were organized into 9 distinct cell types and were also mainly localized to the superficial dorsal...
Figure 3. Neuronal cell types in the human spinal cord
(A) UMAP plot of human spinal neurons showing 35 populations.
(B) Cell2location predictions on spatial transcriptomics data showing selected excitatory (left) and inhibitory (right) cell types at both L3/4 and L5/S1 segmental levels.
(C) Dendrogram showing the relationship of neuronal subtypes, calculated using the top 2,000 highly variable genes and Ward’s method. For each cluster, 2–3 marker genes are listed.

(legend continued on next page)
horn, with the exceptions of Inh-Dorsal-1 and Inh-Dorsal-4 found in lamina III and of Inh-Dorsal-5, which was found in both the superficial and deep dorsal horns (Figures S7 and S9). Expression of the inhibitory transcription factor PAX2, as well as GBX1 and minor expression of LBX1, suggested a dl4/dILA embryonic origin for these cells (Figure S13; Table S3). Most populations expressed markers for both GABAergic (GAD1/2) and glyciner-gic (SLC6A5) cell types, with the exception of Inh-Dorsal-2 and Inh-Dorsal-3, which seemed exclusively GABAergic. The Inh-Dorsal-4 and Inh-Dorsal-5 cell types were putatively non-pepti-dergic and marked by ADARB2, while most other dorsal inhibitory clusters were distinguished by neuropeptide genes such as NPPC, PENK, PDYN, and NPY (Figure S10).

Overall, mid and ventral cells were distinguished by expression of either the early-born markers ZFHx3/4 or the late-born markers NF1A/NFIB/NFIX and PROX1, similar to what has been reported recently in mouse20 (Figure S10). Ex-M-1, Ex-M-2, and Ex-M-3 were mid- excitatory cell types that expressed LMX1B and TLX3, suggesting a dl5/dILB origin (Figure S13). In contrast, Ex-M-4 and the ventral excitatory clusters Ex-V-1, Ex-V-2, and Ex-V-3 expressed very low but detectable levels of markers for the dl1-3 embryonic domains that are known to settle in the deep dorsal and ventral horn regions (BARHL1, BARHL2, LHx2, LHx9, ISL1, and OTP), as well as markers of the V0c (PITX2, CHAT), V2a (VSX2, SOX14), and V3 (SIM1, NKX2-2) domains (Figures S10 and S13). Among inhibitory mid and ventral clusters, all clusters expressed both GABAergic and glycineric marker genes. Inh-M-1 and Inh-M-2 were likely derived from the dl4/dILA domain, based on their expression of either GBX1 or LBX1. Inh-V-2 selectively expressed markers of the V2b embryonic lineage, including GATA2, GATA3, and MSX1. Spatial transcriptomics predictions from Cell2location confirmed a mid/ventral location for each of these clusters (except Ex-M-2) (Figures S7 and S8).

Our characterization of neuronal populations suggested that dorsal clusters were readily distinguished by specific markers, whereas ventral clusters often displayed overlapping patterns of gene expression. Having previously observed this pattern in mouse spinal cord,19 we next systematically assessed how distinct dorsal clusters were compared to ventral clusters in both human and mouse spinal horns. We examined two measures of distinctness: cluster separability (analogous to the tightness of gene expression distribution in a cluster) and distance between cluster centroids (analogous to the difference between means of gene expression).

To assess cluster separability, we used two approaches: silhouette scoring of each cluster (Figures 4A–4C and S11) and a post hoc machine-learning approach to measure the propor-tion of nuclei from each cluster that could be unambiguously assigned (Figure S14A). Both approaches showed that dorsal clusters formed discrete groups that were well separated from each other, whereas mid and ventral clusters were substantially more overlapping (Figures 4B and 4C). To assess the distance between cluster centroids, we calculated pairwise correlations between cluster centroids over the 2,000 most highly variable genes (Figures 4D–4F) and Euclidean distance between cluster centroids in principal-component space (Figure 3C and Russ et al.15). We found that dorsal clusters have significantly lower correlations with each other and higher distances from each other, as compared to mid and ventral clusters, and this was also true for correlations at the individual cell level (Figure S14B).

Overall, these analyses confirmed that dorsal neurons are organized into significantly more discrete groups than ventral neurons, pointing to fundamental differences in the overall organization of the dorsal and ventral regions of both the human and mouse spinal cords.

Having established similarities in the overall organization of dorsal and ventral neurons in the human and mouse, we next examined the extent to which individual gene expression patterns are shared or distinct between the two species. We thus integrated the neuronal data with prior harmonized datasets from postnatal mouse tissue (Figures 5A, 5B, and S15). We found that, overall, human neurons were enriched for KAZN, ROBO2, and DPP10, while mouse neurons were enriched for DCC, USP29, and ASIC2 (Figures 5E and S16). There was cluster-specific correspondence between the two datasets, with pairs of human- mouse dorsal clusters showing high correlations and specific relationships, while ventral clusters showed broader overall similarity (Figure 5Q). We used a network analysis of cluster relat-edness to identify human and mouse cell type pairs with high conservation (Figure 5D). For example, human Ex-Dorsal-4 is highly homologous to mouse Excit-05, a member of the MAF family located in lamina III–IV and associated with corrective reflexes and light touch processing (Figure 5G).21 Both the human and mouse clusters are enriched for MAF, ADARB2, and RORA, while the human cluster is also enriched for MAFA (found in the spatial transcriptomics data in the deeper region of the dorsal horn) and the mechanosensitive protein PIEZO2, which may confer evolutionarily distinct functions in this population. In addition, human Inh-Dorsal-8 was highly homologous to mouse Inhib-11, a member of the Pdyn family located in lamina I–III and associated with mechanical allodynia pain symptoms and itch22–25 (Figure 5L). Both clusters were enriched for the neuropeptides PDYN and PNC, as well as PROX1 and TACR3. The human cluster was enriched for the neuropeptide NPPC, while the mouse cluster was enriched for the neuropeptide Gal. In the future, such cross-species cell type relationships can be used to propose behavioral functions for a broad range of human neuronal populations.

As a resource, this cellular and gene expression atlas of the adult human spinal cord makes it possible to register known molecular correlates of pathophysiology with human spinal cell types. For example, genome-wide association studies (GWAS) have identified a number of human genomic loci...
associated with chronic pain.\textsuperscript{26–28} We examined cell-type-specific expression of the ten genes near loci that were significant in at least two independent cohorts for GWAS analysis (see STAR Methods). Of these, CCDC26 was strongly enriched in microglia and macrophages, while DCC, NOXA1, and SPOCK2 were enriched across the neuronal populations (Figures S16 and S17). Intriguingly, SOX5 was found in astrocytes, vascular cells, and a group of related neuronal cell types (Ex-Dorsal-8 through Ex-Dorsal-12) that are excellent candidates for a role in pathogenic mechanisms of chronic pain. Subsets of these dorsal excitatory peptidergic neurons selectively expressed the TAC1, TAC3, and CALCB genes for pain-related neuropeptides substance P and CGRP; they were enriched for the opioid receptor gene OPRK1, and they selectively expressed the opioid receptor gene OPRD1 (Figure S17). In addition, these cell types were putative homologues of mouse cell types Excit-14 through Excit-19 (Figure 5C), which have been shown to play functional roles in mechanical nociception and pain-coping mechanisms.\textsuperscript{22,29–33} These data link population genetic studies with transcriptional profiling to propose specific cell types as central regulators of chronic pain in human patients.
Human motoneurons are defined by genes related to cell structure, cell size, and ALS

We next sought to use this cellular and molecular resource to study the gene expression profile of human motoneurons and to determine whether their molecular repertoire relates to their selective vulnerability in diseases such as ALS. The human motoneuron cluster could not be divided into more refined discrete subtypes. This may reflect technical limits (these nuclei contained a relatively low number of genes per nucleus) or biological continua among adult human motoneuron features. Co-clustering with mouse motoneurons from previously published datasets suggested a division into alpha/beta and gamma subtypes, but these were not clearly separated by human marker genes (Figures S24A–S24C). As a result, we analyzed human motoneurons as one group.

We examined the top 50 marker genes that distinguished the motoneuron cluster from other human spinal neurons (Figure 6A). To determine whether these genes were enriched in motoneurons in spinal cord tissue, we assessed the distribution of the entire predicted gene signature in our spatial transcriptomics dataset. While a few genes may reflect background contamination (e.g., PLP1), we found that the predicted motoneuron Cell2location distribution and the top motoneuron marker genes were strongly enriched in lamina IX in the ventral horn, confirming the overall expression pattern (Figure S18A). Motoneuron markers included those involved in acetylcholine synthesis and function (SLC5A7 and ACLY), as expected, but surprisingly were dominated by three partially overlapping sets of genes: (1) those involved in cytoskeletal structure, (2) neurofilament genes related to cell size, and (3) genes directly implicated in ALS pathogenesis (Figure 6A).

Cytoskeletal components were the most abundant category of motoneuron marker gene and the most enriched gene ontology (GO) terms, including GO annotation clusters related to microtubules \((p = 0.000009)\) and axon structure and neurofilaments \((p = 0.000018)\) (Table S5). The marker genes that were structural components of neurofilaments (NEFL, NEFM, NEFH, and PRPH) have been directly linked to cell size, axon diameter, and degeneration, providing a potential link between human motoneuron gene expression and cellular phenotype. Among ALS-related motoneuron marker genes, there were both cytoskeletal genes (NEFH, PRPH, TUBA4A, and STMN2) and genes that are not directly linked to cellular structure (SOD1, OPTN, and SPP1). Most of these markers showed enriched expression in lamina IX in the spatial transcriptomics dataset (as examples, Figure S18A).

We further examined the expression of a panel of ALS-related genes compiled from the literature across human spinal cord cell types. In addition to the genes above, CHCHD10 and KIF5A were enriched in spinal motoneurons, extending this signature profile (Figures S18B, S19, and S20). We also observed enriched expression of SPP1, FUS, and C9ORF72 in microglia and STMN2, and TUBA4A in an excitatory mid-population (Ex-M-1, Figures S18B, S19, and S20). TARDBP was not detected at sufficient levels in the dataset to characterize its expression pattern. We next assessed the cell type distribution of co-variying gene modules of recently implicated in ALS pathogenesis based on spatial transcriptomic data. In agreement with our findings above, the main module associated with disease progression in human tissue was enriched in microglia and motoneurons in our dataset (Figure S21).

Given that the enriched expression of neurodegeneration-associated genes in human motoneuron transcriptomics may have been due to the age of the study donors, we also examined expression of ALS-related genes in a dataset of human embryonic spinal cord cell types. We found patterns including low levels of gene expression (i.e., NEFH and TUBA4A), moderate but broad cell type expression (i.e., OPTN and PRPH), or high and ubiquitous cell type expression (i.e., SOD1 and STMN2) (Figure S22). Thus, the overall enrichment of ALS-related genes in human motoneurons was not apparent in newly formed motoneurons but likely emerge at some point during motoneuron maturation or aging. To test whether this expression profile reflected a non-specific enrichment of degeneration-associated genes in human motoneurons with age, we compared the expression of genes for multiple neurodegenerative diseases, including those with age-related associations, across human spinal cord cell types. This analysis revealed a specific association of ALS-related gene expression in human motoneurons (Figure S23).

To determine whether ALS-related genes are also enriched in motoneurons in mice, the major animal model for studying the genetic basis of neurodegenerative disease, we compared the human data to prior snRNA-seq data from lumbar skeletal motoneurons from adult mice. We found that prominent ALS-related genes were enriched and were expressed at higher levels specifically in the human motoneurons as compared with mouse motoneurons (Figures 6B and S24C). To determine whether this enrichment is unique to motoneurons, we examined the analysis of a recent study on conservation in human brain gene expression patterns and found that three genes of interest (SOD1, TUBA4A, OPTN) had a significantly higher mean human-to-mouse divergence score.

Figure 5. Relationships between human and mouse spinal cord neuronal populations

(A and B) UMAP plots of integrated human and mouse spinal neuron datasets, colored by clusters from (A) human dataset and (B) mouse dataset (Russ et al.4). (C) Heatmap of correlation values \((\text{Pearson’s} \; R)\) between human clusters (columns) and mouse clusters (rows). Correlations were calculated using the top 2,000 highly variable genes from the integrated mouse-human dataset. Red boxes highlight 7 pairs of clusters shown in (E–L). Human clusters are bolded, and mouse clusters are in regular font. (D) Quotient graph showing neuronal clusters as nodes connected by edges. Edges represent correlations greater than 0.8 between human and mouse neuronal clusters. Edge thickness and length reflect correlation values, with greater correlations having thicker and shorter edges. Human clusters (bold, pink) and mouse clusters (teal) are shown. Gray circles highlight 7 pairs of clusters shown in (E–L). (E–L) Venn diagrams represent differentially expressed genes using the Wilcoxon rank-sum test. The overlap in the two circles represents top genes enriched in both of the selected pair(s) of human and mouse neurons compared to all other human and mouse neurons, while human or mouse top enriched genes are shown in pink or teal circles, respectively. No differentially expressed genes were found for Mouse Excit-01. See also Figures S15–S17.
than other assayed genes (mean score of 0.587 ± 0.19 versus 1.426 other genes with mean 0.320 ± 0.123, p = 0.0002).

**Cell size and protein expression in human lumbar motoneurons**

Why might human motoneurons be defined by genes related to cell size and structure, compared with other neurons? It is well established that human motoneurons are large, but to further investigate relative size differences, we analyzed neuron soma size across all laminae in human and mouse lumbar spinal cord tissue. Given the obvious differences in overall body size and anatomy, we expected that most classes of human neurons would be larger than mouse neurons. Surprisingly, human and mouse lumbar spinal neurons were approximately the same size, with a median Feret diameter (maximal caliper length) of 16.02 and 13.13 μm, respectively (human mean 20.3 ± 0.28 μm).

Figure 6. Human motoneurons are characterized by genes associated with ALS, cell structure, and increased cell size
(A) Association network plot constructed using the String protein database for the top 50 marker genes of human motoneurons, with selected categories highlighted (cholinergic transmission, orange; ALS, red; genes whose overexpression in mice causes enlargement and/or degeneration of motoneurons, green; cytoskeletal components, gray).
(B) Volcano plot showing genes enriched in either lumbar motoneurons from adult mice or lumbar motoneurons from adult humans. Genes are plotted by the average change in expression (avg. log2-fold change) and by the statistical strength of the difference (−log10(p value)) with significant genes in black and significant ALS-related genes in red.
(C) Gross anatomical and neuronal measurements of the human (H) and mouse (M) lumbar spinal cords, including median neuron size (μm), transverse area of the spinal cord (mm²), maximum nerve length (cm), and body mass (kg).
(D) Transverse sections of one side of the adult lumbar human (above) and mouse (below) spinal cords, with antibody labeling for NeuN. Images are representative of data from three subjects. Scale bars indicate 1 mm. Boxes indicate the regions shown in (E).
(E) Higher magnification view of NeuN-labeled spinal neurons from (D) in the human (above) and mouse (below). The left-side images are from the dorsal horn and the right-side images are of putative motoneurons in lamina IX. Scale bars indicate 125 μm.
(F) Histogram showing the count distribution of neuron Feret distance in human (pink) and mouse (teal) across the different lamina regions of the adult lumbar spinal cord. Measurements are given in μm, and the count scale is shown at the right of each plot. Bonferroni-adjusted Wilcoxon rank-sum test p values and Bhattacharyya Coefficients (BCs) for human versus mouse distributions are as follows. I/II: p = 7.5e−27, BC = 0.93; III/IV: p = 4.0e−12, BC = 0.96; V/VI: p = 3.2e−30, BC = 0.89; VII/VIII: p = 5.7e−49, BC = 0.80; IX: p = 1.6e−19, BC = 0.71; X: p = 9.5e−10, BC = 0.92.
See also Figures S18–S23.
SEM; mouse mean 14.28 ± 0.12 SEM) (Figures 6C and S24D–S24F; Table S6). By contrast, human lamina IX spinal neurons were approximately 2-fold larger than those in mouse and could be up to ∼120 μm across compared with ∼50 μm in mouse (Figures 6D–6F and S24D–S24F; Table S6). These measurements are consistent with those previously reported for human and mouse spinal motoneuron soma [50–52] and the same proportion that has been observed for human and mouse motoneuron axon caliber [53–55]. Assuming that human alpha motoneurons are within the higher end of this size distribution, then they are (1) much larger than other human spinal neurons, (2) increased in scale relative to mouse motoneurons, and (3) among the largest vertebrate neurons, including elephant motoneurons (∼85 μm) [56], human Betz corticospinal neurons (∼60–100 μm) [57], subsets of human dorsal root ganglion neurons (up to 100 μm) [58], and salmon Mauthner cells (∼87 μm) [59]. This notable size of human motoneurons may explain the specialized gene expression signature that we observed in this subclass of neurons.

To assess specific ALS-related features in tissue and in situ cell size, we next analyzed the protein expression of six ALS-related genes in postmortem lumbar spinal cord from four donors using immunofluorescence. We found that neurons expressing NEFH, OPTN, PRPH, STMN2, and TUBA4A proteins were all enriched within the motoneuron region (lamina IX) of the lumbar spinal cord, with limited positive cells in other regions except for scattered, large cells in lamina III/IV (which may be projection neurons) and smaller neurons in medial lamina VII (Figures 7A–7D; Table S7). SOD1 was present in lamina IX and throughout the spinal cord in a distinct peri-nuclear distribution, in contrast to the enriched RNA expression that we detected by snRNA-seq and by spatial transcriptomics. To
ensure the accuracy of the SOD1 protein expression pattern, we validated the SOD1 antibody through targeted knockdown in human induced pluripotent stem (iPS) neurons (Figure S25D). Overall, these data confirm the enriched expression of ALS-related proteins in human spinal motoneurons in tissue. 

We also studied the expression of these proteins in the mouse spinal cord, using lumbar tissue from aged animals (11 months old) to approximate the advanced age of the human subjects in this study. We found that Nefh, Optn, Prph, Stmn2, and Tuba4a displayed enrichment in lamina IX, while Sod1 was expressed ubiquitously, similar to what has been previously described for Sod1 in mice (Figures S25A–S25C). Together with the comparative transcriptomic analysis above, this suggests that while human and mouse motoneurons are both enriched for expression of ALS-related genes, in human motoneurons, the relative RNA expression levels are higher and the enrichment of these genes as motoneuron-specific markers is greater.

Finally, we examined the relationship between expression of ALS-related genes and human spinal neurons size in tissue. We measured the Feret distances of human neurons expressing each ALS-related protein in comparison with non-expressing neurons. We found that neurons that expressed NEFH, OPTN, PRPH, STMN2, and TUBA4A were generally larger than non-expressing neurons, both within the motoneuron region of lamina IX and in other laminae (Figure 7D; Table S7). Within lamina IX, this likely reflects enrichment within the larger alpha motoneurons (versus gamma), and in other laminae, this may reflect expression within spinocerebellar projection neurons that degenerate in ALS.62,63 or other large cell classes. Importantly, we found that the very largest lamina IX neurons—known to be most susceptible to degeneration in ALS62,63—were the most likely to express these markers. For lamina IX neurons with a Feret distance greater than 70 μm, on average, 100% expressed NEFH, 81% expressed OPTN, 88% expressed PRPH, 60% expressed SOD1, 90% expressed STMN2, and 95% expressed TUBA4A (Table S7). These data further link motoneuron size and vulnerability to these cytoskeletal genes that have causative roles in motoneuron size and human disease.

**DISCUSSION**

The advent of single-cell transcriptomic profiling approaches has transformed biology, with the potential to pinpoint therapeutic targets amid the complexity of human disease. However, we still lack a comprehensive characterization of the human spinal cord that could provide crucial insights into chronic pain, spinal cord injury, and neurodegeneration. Here, we used snRNA-seq and spatial transcriptomics to create a cellular atlas of the adult human lumbar spinal cord. We identified dozens of cell types, including diverse glial and neuronal populations, and characterized their molecular repertoires and putative locations. We next used this atlas to examine cell-type-specific mechanisms of pathophysiology, identifying a group of human dorsal horn neurons enriched for pain-related genes, and a specialized molecular signature in human motoneurons that links their extreme cell size with their vulnerability to degeneration in ALS. This atlas and an accompanying web-based resource (https://vmenon.shinyapps.io/humanspinalcord/) can serve as tools for further understanding human spinal cord biology.

There have been several recent studies on the molecular and cellular heterogeneity in the human spinal cord, particularly during development. Rayon and colleagues focused on first trimester spinal cord derived from four human embryos, identified diverse progenitor and neuronal populations, and performed a systematic comparison with the spinal cord cell types of the developing mouse spinal cord. Zhang and colleagues profiled the early and mid-stages of fetal development with an important focus on glial development and cell-cell communication. For the adult human spinal cord, Zhang and colleagues performed snRNA-seq on the spinal cord from two donors and identified coarse glial and neuronal cell types. However, they did not characterize human neurons to the same degree as this study, especially with respect to motoneurons, nor did they validate predicted gene expression patterns in tissue.

Here, we establish the first comprehensive taxonomy of the adult human spinal cord. With this broad view and comparison with similar work in mice, we found that the major axes of spinal neuron diversity are conserved across both species. In addition to neurotransmitter status, the primary factor in spinal neuron organization is dorsal-ventral location, with dorsal neurons forming robust and distinct clusters that each display specific marker genes while ventral clusters showed overlapping gene expression patterns. This is similar to what we and others have previously shown in adult mice, but seems to be at odds with our knowledge of the many refined populations of ventral neurons within the cardinal V0, V1, V2, and V3 embryonic lineage domains. There are multiple potential explanations for this discrepancy. First, it is possible that the dorsal-ventral pattern reflects multiple axes of neural diversity overlaid onto ventral neurons, effectively blurring the distinctions that would be apparent along any single axis. In addition to developmental lineage, it has recently been shown that transcriptional signatures of birthdate sub-divide spinal neurons during mouse and human development. This includes a continuum of very early-born (ONECUT2), early-born (ZFHX3/4), and late-born (NFIB/NFIA/NEUROD2/6) factors and is most apparent among cell types that settle in the mid and ventral regions of the spinal cord. As birthdate is coupled to projection neuron versus local interneuron identity, this level of transcriptional diversity may be sustained into adult stages to support cell-type-specific functional requirements based on axon length. Together with other parameters, such as location within the ventral horn or electrophysiological specialization, these features may overlap each other to form broad ventral clusters that are less distinct from each other. Alternatively, it is possible that dorsal and ventral neurons in the adult require differential levels of ongoing gene expression related to their functions. Perhaps dorsal neurons (which are marked by specific neuropeptides, neuropeptide receptors, and other genes involved in neuronal function) require specific transcriptional signatures to be sustained to perform more specialized computations. In contrast, ventral neurons may operate mainly based on their connectivity within the network and can therefore downregulate their lineage-based and embryonic molecular diversity once axon guidance has occurred and circuit structure is complete. Relatedly, ventral
neurons may operate as a broad network whose tasks are carried out through global dynamic population activity. In the future, relating transcriptional identity to connectivity, intrinsic electrophysiological parameters, and neural activity will help to resolve how spinal neural populations are organized for function.

An intriguing finding from our analysis is the enrichment of cytoskeletal gene expression in human motoneurons. All cells require a functional cytoskeleton, raising the question of why spinal motoneurons are particularly dependent on the proper expression and function of cytoskeletal-related genes. Interestingly, neurofilament genes that were enriched in human spinal motoneurons are precisely those structural components that drive increased axon caliber and cell size. Overexpression of mouse NEFL, human NEFM, human NEFH, or mouse PRPH in transgenic mice can each cause enlargement and swellings of motoneuron somas and subsequent axon degeneration, linking human motoneuron gene expression and potential degenerative phenotypes. Relatedly, these neurofilament genes are found in other large neurons in the nervous system, suggesting that they may be part of a common signature that permits increased cell size. Large soma size and axon caliber may be required to sustain extensive dendritic trees and axons up to a meter long, to support cell energetics, or for firing rate and conduction parameters. These large cells then rely critically on this protein network and are selectively vulnerable to its abnormal function. Human motoneurons were also distinguished by expression of the microtubule stability factors TUBA4A and STMN2, potentially highlighting a requirement for structural support in these peripherally projecting cells with long axons. Overall, these findings support a model of specific molecular repertoires for motoneuron cell structure that also confer selective vulnerability to degeneration.

Although we captured all major cell types and most known subclasses of cells in this work, we foresee further advances as additional datasets of this type arise. Among motoneurons, we expected to observe well-established “alpha” and “gamma” subtypes based on transcriptional profiles recently described in mice but did not. This limitation may be experimental, reflecting the relatively low number of genes detected per motoneuron nucleus. In addition, we did not observe all known populations of neurons, such as cerebrospinal contacting neurons, which represent less than 1% of mouse spinal neurons. As technological advances allow for higher-sensitivity transcriptomics on larger numbers of cells or improvements in in situ profiling methods, a clearer picture of refined neuronal populations and the heterogeneity within motoneurons will likely become apparent.

Overall, it is important to consider the spinal cord as a community of cell types that function together in normal health and disease. Here, we highlighted specific findings on a proliferative population of adult microglia, a group of dorsal excitatory neurons enriched for pain-related genes, and the molecular signature of motoneurons. As a whole, this work provides a comprehensive resource for transcriptional profiling of the dozens of cell types that make up the adult human lumbar spinal cord. As such, it will allow researchers to parse how genetic alterations could affect diverse cell-type-specific molecular profiles in disease; how particular populations may respond to target molecular interventions and pharmacology; and how human spinal cell types may interact with each other through cell-cell signaling pathways. Thus, we hope this work, together with other ongoing efforts, will serve as a foundation for studying the wide range of cell types involved in human spinal cord function.

STAR METHODS

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SUPPLEMENTAL INFORMATION

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DECLARATION OF INTERESTS

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INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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## Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| KI67                | Cell Signaling Tech | 9449S |
| IBA1                | Synaptic Systems | 234006 |
| NeuN                | Millipore Sigma | ABN90P |
| SOX9                | Abcam | ab185966 |
| OLIG2               | Millipore Sigma | MABN50 |
| SOD1                | Sigma | HPA001401 |
| OPTN                | Proteintech | 10837-1-AP |
| Neurofilament H     | Cell Signaling | 2836S |
| Chat                | Millipore Sigma | AB144P |
| TUBA4A              | ThermoFisher | PAS-29546 |
| Alexa Fluor® 647 Anti-alpha Tubulin | Abcam | ab190573 |
| Stathmin-2/STMN2    | Novus | NBP1-49461 |
| Peripherin/PRPH     | Millipore | AB1530 |
| **Biological samples** |       |            |
| Human spinal cord   | Ottawa Hospital Research Institute | N/A |
| Human spinal cord   | Gui de Chauliac Hospital | N/A |
| Human spinal cord   | Target ALS Multicenter Post-mortem Core | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| ViralBoost Reagent  | ALSTEM | #VB100 |
| Lenti-X concentrator | Takara Bio | #631231 |
| Matrigel            | Corning Incorporated | #354277 |
| Hoechst             | Thermo Scientific | #62249 |
| Pro-Long Gold antifade reagent | Invitrogen | #P36934 |
| Sucrose             | Invitrogen | 15503-022 |
| 1 M HEPES (pH = 8.0) | Gibco | 15630-080 |
| CaCl2               | Sigma Aldrich | C1016-100G |
| MgAc                | Sigma Aldrich | M5661-50G |
| 0.5 M EDTA (pH = 8.0) | Corning | MT-46034Cl |
| Dithiothreitol (DTT) | Sigma Aldrich | 10197777001 |
| Triton X-           | Sigma Aldrich | T8787 |
| 1 M Tris-HCl (pH = 7.4) | Sigma Aldrich | T2194 |
| 0.04% BSA           | New England Biolabs | B9000S |
| 0.2 U/μL RNase Inhibitor | Lucigen | 30281-1 |
| Trypan Blue Stain (0.4%) | Thermo Fisher Scientific | T10282 |
| **Critical commercial assays** | | |
| Chromium Single Cell 3' GEM, Library & Gel Bead Kit V3 | 10X Genomics | PN-1000075 |
| Chromium Single Cell B Chip Kit | 10X Genomics | PN-1000074 |
| Chromium i7 Multiplex Kit | 10X Genomics | PN-120262 |
| Visium Spatial Gene Expression Slide & Reagent Kit | 10X Genomics | PN-1000184 |
| Visium Accessory Kit | 10X Genomics | PN-1000194 |
| Dual Index Kit TT Set A | 10X Genomics | PN-1000215 |

(Continued on next page)
### REAGENT or RESOURCE SOURCE IDENTIFIER

**Deposited data**

| Deposited data                      | This paper | GEO: GSE190442 and GSE222322 |

**Experimental models: Cell lines**

| Human iPSC Line | Tian et al. [82] | N/A |

**Experimental models: Organisms/strains**

| C57BL/6J         | Jackson Laboratory | 000664 |
| BALB/cJ          | Jackson Laboratory | 000651 |

**Recombinant DNA**

| mU6-sgRNA EF1a-puro-T2A-2XmycNLS-BFP | Tian et al. [82] | Addgene Plasmid pMK1334 #127965 |

**Software and algorithms**

| Seurat 4.0       | Hafemeister and Satija [83], Stuart et al. [84] | https://satijalab.org/seurat/index.html |
| biomaRt          | Durinck et al. [85] | https://bioconductor.org/packages/release/bioc/html/biomaRt.html |
| Homologene       | National Center for Biotechnology Information | https://CRAN.R-project.org/package=homologene |
| Neurodegenerative disease gene analysis | This paper | https://colab.research.google.com/drive/19Ty97LOwT3AmVCJGKA8BXN FYH_iSZ4?usp=sharing and https://colab.research.google.com/drive/1BDJaixVhYnhMO9VJZNWUn9lw_c87Y7mr?usp=sharing |
| Cell2Location    | Kleshchevnikov et al. [86] | https://cell2location.readthedocs.io/en/latest/ |
| OpenCV           |opencv-python-headless 4.6.0.66 | https://pypli.org/project/opencv-python-headless/ |
| Fiji/ImageJ      | Fijiv2.1.0 | https://imagej.net/software/fiji/ |
| Adobe Photoshop  | Adobe Systems | https://www.adobe.com |
| Adobe Illustrator| Adobe Systems | https://www.adobe.com |
| Other            | This paper | https://vmenon.shinyapps.io/humanspinalcord/ |

**RESOURCE AVAILABILITY**

**Lead contact**

Further information and requests for resources, reagents, or code should be directed to and will be fulfilled by the lead contact Ariel Levine (ariel.levine@nih.gov).

**Materials availability**

The study did not generate new unique reagents.

**Data and code availability**

Anonymized raw sequencing data and counts tables are deposited in the Gene Expression Omnibus (GEO) with accession numbers GEO: GSE190442 and GEO: GSE222322 with associated metadata in Table S2. In addition, visualization of expression data at the cluster and donor level are available through a searchable web resource at https://vmenon.shinyapps.io/humanspinalcord/.

Code for the evolutionary divergence analysis can be found at https://colab.research.google.com/drive/19Ty97LOwT3AmVCJGKA8BXN FYH_iSZ4?usp=sharing and https://colab.research.google.com/drive/1BDJaixVhYnhMO9VJZNWUn9lw_c87Y7mr?usp=sharing. Custom MATLAB-based code for quantification of cell counts is available at https://github.com/ArielLevineLabNINDS/CellCounter (https://doi.org/10.5281/zenodo.6967482).
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Lumbar spinal cords were obtained from neurologic determination of death organ-donor patients (all demographic data are listed in Table S1) under the approval of the French institution for organ transplantation (Agence de la Biomédecine) or the Ottawa Health Science Network Research Ethics Board, following the template provided by the University of Ottawa and the Tri-Council Policy Statement Guidelines. Both approvals imply consent for using anonymized donor genetic information. Patients with neurological disease or major infections were excluded from the study.

For mouse experiments, all mice were of 50:50 mixed background from strains C57BL/6J and BALB/CJ, housed in standard conditions. For basic anatomical experiments, two male and two female mice of approximately 24 weeks old were used. For ALS marker gene expression studies, two male and two female mice of approximately 11 months old were used. All procedures and experiments were approved by the Animal Care and Use Committee of NINDS (protocol #1384).

METHOD DETAILS

Human spinal cord acquisition and preparation

Human lumbar spinal cords were retrieved under chilled body and neuroprotective conditions as described previously.87–89 The extraction procedure took 20–40 min and was done within 3 h of cessation of circulation by aortic cross-clamp. For single nucleus RNA sequencing experiments, lumbar spinal cord tissue from donors (~50–80 years old, 4 men, 3 women) was flash frozen on liquid nitrogen in the operating room and stored at −80 °C until nuclei isolation.

For immunohistochemistry experiments, lumbar spinal cord tissue was isolated from organ-donor patients (~55–65 years old, 3 men, 1 woman). The tissue was immediately fixed in 4% paraformaldehyde for 24–48 h, then washed in PBS, and placed in 30% sucrose for 2–4 days at 4 °C before being embedded in OCT medium for sectioning.

For Visium spatial transcriptomics, post-mortem lumbar spinal cord from a non-neurological control subject (~75 years old, male) was acquired from the Target ALS Multicenter Post-mortem Core as part of the New York Genome Center (NYGC) Amyotrophic Lateral Sclerosis (ALS) Consortium. Informed consent is acquired by each Target ALS member site through its own institutional review board (IRB) protocol and samples are transferred to the NYGC in accordance with all applicable foreign, domestic, federal, state, and local laws and regulations for processing, sequencing and analysis. The Biomedical Research Alliance of New York (BRANY) IRB serves as the central ethics oversight body for the NYGC ALS Consortium. Ethical approval for this study was given by the BRANY IRB.

Mouse work and spinal cord acquisition

All procedures and experiments were approved by the Animal Care and Use Committee of NINDS (protocol #1384). Adult mice were of 50:50 mixed background from strains C57BL/6J and BALB/CJ, housed in standard conditions. For basic anatomical experiments, two male and two female mice of approximately 24 weeks old were used. For ALS marker gene expression studies, two male and one female mice of approximately 11 months old were used. To obtain spinal cord tissue, anesthetized mice were transcardially perfused with PBS followed by cold 4% paraformaldehyde (PFA). The spinal cords were harvested and post-fixed in 30% sucrose overnight at 4 °C, cryoprotected by immersion in 30% sucrose overnight at 4 °C and embedded in OCT medium for sectioning.

Nuclei isolation

Nuclei were isolated from fresh frozen human spinal cords using a triton-based protocol.90 Briefly, after removing the dura, half a segment of spinal cord was placed in a Dounce homogenizer (Kontes Dounce Tissue Grinder) containing 500 μL of lysis buffer (0.32 M sucrose, 10 mM HEPES [pH 8.0], 5 mM CaCl2, 3 mM MgAc, 0.1 mM EDTA, 1 mM DTT, 0.1% Triton X-100). After douncing with 5 strokes of pestle A and 5–10 strokes of pestle B, the lysate was diluted in 3 mL of sucrose buffer (0.32 M sucrose, 10 mM HEPES [pH 8.0], 5 mM CaCl2, 3 mM MgAc, 0.1 mM EDTA, 1 mM DTT) and passed over a 70 μm strainer. The filtered lysate was centrifuged at 3,200 x g for 5 min at 4 °C. After centrifugation, the pellet was resuspended in 3 mL of sucrose buffer and centrifuged again at 3,200 x g for 5 min at 4 °C. After centrifugation, the pellet was resuspended in 3 mL sucrose buffer and incubated for 2 min on ice. The sample was transferred to an Oak Ridge tube and homogenized for 1 min using an Ultra-Turrax Homogenizer (IKA). Then, 12.5 mL of density sucrose buffer (1 M sucrose, 10 mM HEPES [pH 8.0], 3 mM MgAc, 1 mM DTT) was layered below the sample. The tube was centrifuged at 3,200 x g for 20 min and the supernatant immediately poured off. The nuclei on the side of the tube were resuspended with 100 μL of PBS with 0.04% BSA and 0.2 U/μL RNase inhibitor. Nuclei were inspected for visual appearance and quantified with a hemocytometer before proceeding with nuclei capture and sequencing.

Single-nucleus RNA sequencing

Single nucleus RNA sequencing was carried out using Single-cell gene expression 3’ v3 kit on the Chromium platform (10X Genomics) according to manufacturer’s instructions with one modification. Following reverse-transcription, an additional PCR cycle was added to the number of cycles for cDNA amplification to compensate for decreased cDNA abundance in nuclei compared to
cells. Libraries were sequenced to a minimum depth of 20,000 reads per nucleus using an Illumina HiSeq 3000 (PE 26–8 – 98 bp). Raw sequencing reads were demultiplexed, aligned, and a count matrix was generated using CellRanger. For alignment, introns and exons were included in the reference genome (GRCh38).

**Quality check analysis**
All the 10x runs for each human sample were initially filtered with an nUMI cutoff of >1000 and then nuclei with less than 5% mitochondrial gene contamination were retained. Next, the mitochondrial genes were also removed from the matrices. A total of 55,289 nuclei that passed quality control filtering, with mean detection of 2,187 genes per nucleus (Figure S1).

**Top level UMAP and clustering**
The 7 human datasets were integrated using SCTransform normalization followed by CCA based integration using the Seurat 4.0 package. The integrated datasets were then jointly analyzed to identify the optimal number of principal components for downstream analysis, based on the ElbowPlot and PCheatmaps function. The number of PCs was set to 30 for clustering and UMAP visualization. The clusters, obtained using a value of 0.6 for Seurat’s resolution parameter, were then manually annotated based on the expression of marker genes for neurons, astrocytes, microglia, oligodendrocytes, OPCs, endothelial cells, pericytes, meningeal cells, Schwann cells, and lymphocytes.

**Subclustering of major cell types**
Identification of sub-clusters within cell types was performed separately for three major cell types (neurons, microglia, astrocytes), with the rest being sub-clustered in two additional groups (Group 1- oligodendrocytes, OPCs, and Schwann cells; Group 2- endothelial cells, pericytes, meningeal cells, and lymphocytes). For each cell type/group, the subclustering was done in multiple rounds until no putative transcriptomic doublets or contamination of other cell types was observed, as described below.

For subclustering of major cell types, the raw counts were aggregated from all 7 datasets for each cell type, and then re-normalized (using log normalization) and scaled in order to prepare for integration. The integration of 7 datasets belonging to a particular cell type was performed based on the CCA-integration workflow from the Seurat 4.0 package. The optimal number of PCs was selected based on the ElbowPlot and PCheatmaps function for each cell type, in order to be used for subclustering and preparation of UMAP visualizations. Multiple resolutions were interrogated, depending on cell type, ranging from values of 0.08–3.

During each round, putative transcriptomic doublet clusters and contamination of other cell types was removed (based on co-expression of multiple major class genes) and the above steps were performed again. Doublets were identified by clusters that expressed markers for more than one cell type. All clusters were checked for doublets by their markers using the wilcox (Wilcoxon Rank-Sum Test) and auroc (Area under the ROC curve) functions, as well as visually using the FeatureScatter option in Seurat.

**Subclustering of neurons**
Neurons were clustered in 2 stages. During the first stage, log-normalization of raw counts and scaling (including regressing out the number of transcripts and the percentage of mitochondrial transcripts) of each dataset were performed, followed by integration based on the same steps described above. All high-quality neuronal nuclei were clustered in an unsupervised manner testing a range of resolutions. This stage led to some clusters that were clearly distinct, together with a large group of nuclei (visualized in the central region of the UMAP) that did not sub-divide well at lower resolutions, but were dispersed at higher resolutions. This prompted us to perform a focused re-clustering of this non-distinct group with new principal components based only on these nuclei. The spatial distribution of marker genes for this large group suggested a ventral identity, so we included all putative mid/ventral clusters into this step of the analysis. Raw counts were again extracted and normalized using SCTransform (to avoid dataset size related limitations), followed by the standard integration workflow in the Seurat 4.0 package. We also performed a focused subclustering of the motoneurons at this stage but could not identify robust sub-clusters within this cell type. During each clustering stage, cluster-specific genes were identified based on Wilcoxon Rank-Sum test and AUROC analysis within the FindMarkers function from Seurat 4.0. Based on these genes, distinct subpopulations based on expression of candidate markers were manually annotated.

In order to obtain a refined set of neuronal subpopulations, all the sub-clusters were interrogated for low gene detection, doublets, and other contamination from non-neuronal genes, and were subsequently removed from the analysis. All the refined clusters were then re-integrated to prepare a combined neuronal UMAP and mapped with refined subcluster annotations. Finally, we then analyzed each cluster independently to assign a “dorsal”, “mid”, “ventral” identity, guided by spatial distribution of marker genes, the predicted cell2location distribution of the cell type in spatial transcriptomics data, and relative similarity to previously described mouse or macaque cell types.

**Cluster robustness assessment and silhouette scores**
We used two approaches to assess cluster robustness: a post-hoc machine learning-based classification approach, and a silhouette score approach.

For the post-hoc machine learning approach, we built a random forest classifier for every pair of neuronal clusters, trained on 80% of the nuclei. This classifier was then used to assign cluster membership for the remaining 20% of the cells, and the entire process repeated such that each cell in every pairwise cluster comparison was classified 100 times. A cell that was classified into its original
cluster <90 times was deemed “misclassified”. For every pair of clusters, we then calculated the mean percentage of cells that were misclassified among the two clusters to generate pairwise cluster robustness scores. For visualization as a constellation diagram, we only connected cluster pairs with minimum misclassification percentage >3%, representing their connections with the mean misclassification percentage.

For silhouette score evaluation, we used the ‘silhouette’ function from the ‘cluster’ library in R (https://cran.r-project.org/web/packages/cluster/index.html), where the Euclidian distance matrix based on the first 25 PCs was used as input, together with the neuronal cell type annotations.

**Tissue processing, Visium data generation, and Visium data preprocessing**

Frozen post-mortem lumbar spinal cord from 5 non-neurological control subjects were embedded in Tissue Plus OCT Compound (Fisher Healthcare, catalog no. 4585) and cryosectioned at –16°C. Sections of 10 μm thickness were collected onto prechilled Visium Spatial Gene Expression Slides (10x Genomics, catalog no. 1000185) by warming the back of the slide to adhere the tissue. Four technical replicates for each subject were collected approximately 30 mm apart across two Visium slides that were processed in parallel to minimize technical batch effects. Where necessary due to tissue sectioning artifacts, additional technical replicates were collected and processed on a later date. All technical replicates for each subject were sequenced on the same flow cell.

Visium spatially resolved gene expression data were generated according to the Visium Spatial Gene Expression User Guide (10x Genomics, CG000239 Rev F). Briefly, tissue sections were fixed in chilled methanol and stained using hematoxylin and eosin. Brightfield histological images were acquired using an EC Plan-Neofluar 10x/0.3 M27 objective on a Zeiss Axio Observer Z1 fitted with a Zeiss Axioscam 506 mono (Carl Zeiss Microscopy, Germany). Raw CZI images were stitched using Zen 2012 (blue edition) (Carl Zeiss Microscopy, Germany) and exported as JPEGs. Tissue sections were permeabilized for 12 min which was selected as the optimal time based on tissue permeabilization time course experiments conducted using the Visium tissue optimization protocol. cDNA libraries were prepared and quantified according to the Visium Spatial Gene Expression User Guide (10x Genomics, CG000239 Rev F) and pooled at a concentration of 10 nM for sequencing.

Pooled spatial gene expression libraries were loaded at a concentration of 0.9 nM and sequenced on a NovaSeq 6000 System using a NovaSeq S4 Regent Kit v1.5 (200 cycles, Illumina, catalog no. 20027466) using the following recipe: read 1: 100 reads, i7 index read: 10 cycles, i5 index read: 10 cycles, read 2: 100 cycles. The average sequencing depth for each sample was approximately 200-280 x 106 reads.

Raw FASTQ files and histological images were processed using Space Ranger v.1.3.0, which uses a modified STAR v2.7.2a for genome alignment and performs barcode/UMI counting to generate feature-spot matrices. Reads were aligned to a GRCh38 reference genome filtered to exclude IncRNAs, pseudogenes and mitochondrially encoded genes.

**Postprocessing and computational analysis of Visium spatial transcriptomics data**

Registration of Visium experiments was achieved using the OpenCV library (https://pypi.org/project/opencv-python-headless/). Briefly, histology images were annotated with 31 landmarks (14 points denoting features of the gray-white matter boundary for each side, and 3 points along the midline). These points were used to calculate a homography matrix from each experiment to a reference image using the findHomography function. Coordinates from each experiment were transformed to the resulting coordinate space using the perspectiveTransform function.

Cell type proportions comprising Visium experiments were estimated using the Cell2location package based on the following notebook (https://cell2location.readthedocs.io/en/latest/notebooks/cell2location_tutorial.html). The single-cell regression model was trained with the following parameters: max_epochs = 250 and lr = 0.002. The Cell2location model was initialized and trained with the following parameters: N_cells_per_location = 5, detection_alpha = 20, and max_epochs = 2000.

**Cross-species analysis between human spinal cord vs mouse meta-analysis datasets**

Cross-species comparison between human and mouse meta-analysis spinal cord datasets were performed at two levels: 1. “Top-level”, which includes all major cell types and 2. Neurons only.

In both cases, the orthologous genes within mouse data matrix were converted to human homologs using biomartR package from Bioconductor and in-house scripts. The raw counts from both human and mouse datasets were then split by different samples and then re-normalized, scaled and integrated. For the “top-level” analysis, SCTransform based integration was performed whereas for neurons only, log normalization-based integration was performed. Subsequently, UMAPs and correlation matrices were generated for further cross-species comparison of various cell types at top level and neuronal sub-clusters.

**Cross-correlation of human and mouse cluster expression**

Cross-species cluster correlation measures were calculated from PCs in the integrated space (using 20 PCs for the top-level comparison of major cell classes), and Pearson correlation of the top 2,000 highly variable genes. Aggregate correlation values for each pair of clusters (one mouse, one human) were calculated as the mean correlation value across all human-mouse nuclei pairs from the respective clusters.

Quotient graphs using qgraph in R were used to show the correlations greater than 0.8 based on the top 2,000 highly variable genes between human and mouse spinal cord neurons (graph “cor”, layout “spring”).
GO analysis of human motoneuron marker genes

The top markers (based on smallest adjusted p value) of human spinal motoneurons were determined based on the Wilcoxon Rank-Sum test and were analyzed using DAVID 6.8 GO enrichment analysis (https://david.ncifcrf.gov/summary.jsp). The general categories of GOTERM_BP_DIRECT, GOTERM_CC_DIRECT, and GOTERM_MF_DIRECT were analyzed and functional annotation clustering was performed using default parameters including medium classification stringency.

Focused comparison of mouse and human motor neurons

Human motor neurons were compared to mouse lumbar skeletal motor neurons from a recent study. Mouse MN genes were converted to human homologs using Homologene (https://CRAN.R-project.org/package=homologene). Only genes with human homologs present in both datasets were included in the analysis (13,574). Raw counts were extracted from each original dataset, normalized using SCTransform, and integrated based on integration anchors. Clustering was performed as described above (resolution = 0.4), and differentially expressed genes were identified based on Wilcoxon Rank-Sum test and ROC analysis within FindMarkers function from Seurat 4.0.

Analysis of evolutionarily convergence/divergence scores

All available data on gene expression-based human:mouse divergence scores were downloaded from Pembroke et al. Genes of interest were then extracted, yielding scores for three genes (SOD1, TUBA4A, OPTN) that overlapped with these data. We compared the mean and SD of these three genes to the same metrics for the remainder of the assayed genes from the Pembroke report (N = 1426 other genes) using a standard two-sided t test.

Pain GWAS analysis

All genes found in both Johnston et al. and Suri et al., or in Bortsov et al. (which included separate discovery and replication cohorts) were examined: C8orf34, CCDC26, DCC, EXD3, MIPOL1, NOXA1, PSAP, SLC25A21, SOX9, SPOCK2.

Neurodegenerative disease gene analysis

Post-QC scRNAseq count data was extracted for seven major cell classes of interest. For each gene per cell class, mean expression was calculated across all assayed cells of that class. These means were then z-scored to facilitate comparisons across multiple cell types. Additionally, genes that did not have any count-based data available for that cell class were set to zero for z-scaled values. From this large dataframe of normalized counts per cell type, candidate genes for HSP, PD and ALS were extracted from Genomics England Expert Panel App genes audited at the "green" level of confidence (https://panelapp.genomicsengland.co.uk/). AD genes were annotated by an expert panel and extracted and the ALS list was also supplemented with genes from the literature, as described in the main text. These extracted genes were then mapped to each cluster using the python package “seaborn”, with z-scores greater than 7 truncated to a value of 7 for display purposes. Code for the evolutionary divergence analysis can be found at https://colab.research.google.com/drive/19Ty97LOwT3AmaVCJGKAwBYNXFyH_ISZ4?usp=sharing and https://colab.research.google.com/drive/1BDJaiwhYnhMO9VJZNWUn9lw_c87Y7mjr?usp=sharing.

SOD1 antibody validation in human iPS neurons with targeted knockdown

Previously published human inducible pluripotent stem cells (hiPSCs) were used to knock down SOD1. A SOD1 or non-targeting control sgRNA was cloned into an mU6-sgRNA EF1a-puro-T2A-2XmycNLS-BFP vector (gift from Martin Kampmann’s lab; Addgene #127965). sgRNA sequences are as follows: SOD1: GAGGCACCACGACAGACCCG, non-targeting sgRNA: GAATATGTGC GTGCATGAAG. Lentivirus was produced via transduction of Lenti-X HEK 293T cells using Lipofectamine 3000 in DMEM high glucose GlutaMax Supplement media containing 10% FBS. 24 h post-transfection, media was replaced, including ViralBoost Reagent (ALSTEM, #VB100). 96 h post-transfection, media was collected and concentrated 1:10 in 1xPBS using Lenti-X concentrator (Takara Bio, #631231), aliquoted, and stored at −80°C. 100 mL of these aliquots was used to transduce 100,000 hiPSCs to generate SOD1 KD and control lines. The cells were split and replated on Matrigel (Corning Incorporated #354277) coated coverslips with the viral concentrate in E8+Y-27632 ROCK Inhibitor and allowed to incubate for 24 h at 37°C, 5% CO2. The media was replaced with E8 and the cells were allowed to grow for another 24 h before fixation with 4% PFA in PBS for 10 min at room temperature. Cells were washed with PBS 3 times and permeabilized in block (PBS +3% donkey serum +0.1% Triton X-) for 30 min at room temperature. Primary antibody targeting SOD1 (Sigma, HPA001401-100UL) was diluted at 1:500 in block and cells were incubated in primary overnight at 4°C on a rocker. The next day, cells were washed three times with PBST and incubated in block with secondary antibody (Jackson ImmunoResearch # 711-625-152) and Hoechst (Thermo Scientific #62249) for 1 h at room temperature. Following 3 washes with PBST, the coverslips were mounted using Pro-Long Gold antifade reagent (Invitrogen #P36934). After curing, the coverslips were imaged using Nikon spinning disk confocal using laser wavelengths of 405 nm, 488 nm, 561 nm, and 640 nm at 100 ms exposure and 75%, 25% and 100% power respectively. Images were edited using ImageJ.

Immunohistochemistry antibodies

K67 (Cell Signaling Tech, 9449S), IBA1 (Synaptic Systems, 234,006), NeuN (Millipore Sigma, ABN90), SOX9 (Abcam, ab185966), OLIG2 (Millipore Sigma, MABN50), SOD1 (Sigma, HPA001401-100UL), OPTN (Proteintech, 10837-1-AP), Neurofilament H (Cell

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NeuroResource
Immunohistochemistry

Immunohistochemistry for human and mouse spinal cords were performed as previously described with modifications for human spinal cords.

For single-round immunohistochemistry, human spinal cords were cut at 14 μm, washed twice in TBS and placed in 0.05% sodium azide-TBS at 4°C for 3 days under an LED light to quench autofluorescence. This was done to quench the background autofluorescence and was based on a protocol optimized in brain tissue. Human spinal cords were then placed in blocking buffer (1% IgG-free BSA, 10% normal donkey serum, in TBS) for 1 h prior to incubation in blocking buffer and primary antibody for 48 h at 4°C. Primary antibody was washed off three times in TBS with 0.025% triton before a 2-h incubation in secondary antibody at room temperature. Secondary antibody was washed off three times in TBS with 0.025% triton before adding a coverslip. For multiplex immunohistochemistry, human spinal cords were cut at 10-μm and tissue was processed as previously described.

We examined whether the human tissue underwent size changes during processing that could bias cell size measurements, but we found that processed and stained sections showed overall dimensions that were very similar to the cross-sectional dimensions of lumbar spinal cord from in vivo human MRI. Specifically, in sections from four donors over L3-L5, we measured 9.2 ± 0.17 cm in width and 6.9 ± 0.1 cm in height (mean ± SEM), while Toossi et al. showed that the L3-L5 spinal segments in MRI were ~8.5-9.3 cm in width and 7-7.5 cm in height and, in sections from four donors over L3-L5, we measured 9.2 ± 0.17 cm in width and 6.9 ± 0.1 cm in height (mean ± SEM).

Mouse spinal cords were cut at 50 μm and placed in blocking buffer (1% IgG-free BSA, 10% normal donkey serum, 0.1% Triton X-100 in PBS) for 1 h, then incubated in blocking buffer with primary antibody for 48 h at 4°C. Primary antibody was washed off three times in PBS before a 2-h incubation in secondary antibody at room temperature. Secondary antibody was washed off three times in PBS before adding a coverslip. There was less autofluorescence in mouse spinal cord tissue compared to human spinal cord tissue, giving the ability to perfuse with PBS and PFA, timing of euthanasia and age of the mice. This rendered an LED quenching step unnecessary in mouse tissue.

Imaging

Images of immunohistochemistry samples were imaged using a Zeiss 800 LSM confocal microscope.

Image analysis and quantification

For quantification of neurons by laminae, the images were overlaid in Adobe Photoshop where borders between the gray and white matter and the lamina within the gray matter were drawn. These images were then exported to ImageJ for analysis. The cells were measured manually by outlining each cell using the selection tool and adding them to groups within the ROIManager in ImageJ based on lamina. Feret distance (maximum caliper, similar to diameter) measurements of all the ROIs for each section were saved in a spreadsheet. The white and gray matter of each subject were outlined in ImageJ and their areas were exported to a spreadsheet.

To identify colocalization of markers with NeuN, each neuron was first outlined with the selection tool in ImageJ and saved into different groups based on whether the cell was in lamina IX or not. Then, each cell that had co-occurrence of the markers were placed into separate groups (double-positive in lamina IX and double-positive outside lamina IX). Feret diameter measurements were then saved to a spreadsheet and the number of cells in each group were counted in Python.

For neuron and non-neuronal quantification in human tissue, all counts were done on single thin sections from each donor (10 μm). DAPI was counted using a custom MATLAB-based code and only cells in which the nucleus (DAPI+) was present in the section were subsequently counted for antibody staining (NeuN, Oligo2, Iba1 and Sox9) which was done using the Count Tool in Adobe Photoshop. We ensured that cells on stitching boundaries were not double or miscounted. While a low level of background of antibody staining was present in some cases, we counted only cells which had a clear staining without high background.

QUANTIFICATION AND STATISTICAL ANALYSIS

Two-way ANOVA (repeated-measures) was used for assessing grouped data, such as the correlation and silhouette scores between human and mouse dorsal vs ventral neurons. Two-tailed t tests (unpaired) were used all for differences in silhouette scores and correlation between clusters as well as expression of protein and soma size, as indicated in figure legends. Bonferroni-adjusted Wilcoxon Rank-Sum test p values and Bhattacharyya Coefficients (BC) were used for comparison of human vs mouse cell diameter. Differences among groups were considered significant if p < 0.05. p values are denoted by asterisks: *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001; n.s – not significant. Data are represented as mean ± SEM unless otherwise indicated. Statistical analyses were performed using GraphPad prism software and R.