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11	Mapping of the podocin proximity-dependent proteome reveals novels components of the
12	kidney podocyte foot process
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#### 50 Abstract

The unique architecture of glomerular podocytes is integral to kidney filtration. Interdigitating foot processes extend from the podocyte cell body, wrap around fenestrated capillaries, and form specialized junctional complexes termed slit diaphragms to create a molecular sieve. However, the full complement of proteins which maintain foot process integrity, and how this localized proteome changes with disease, remains to be elucidated. Proximity-dependent biotin identification (BioID) enables the identification of spatially localized proteomes. To this end, we developed a novel in vivo BioID knock-in mouse model. We utilized the slit diaphragm protein podocin (Nphs2) to create a podocin-BioID fusion. Podocin-BioID localizes to the slit diaphragm and biotin injection leads to podocyte-specific protein biotinylation. We isolated the biotinylated proteins and performed mass spectrometry to identify proximal interactors. Gene ontology analysis of 54 proteins specifically enriched in our podocin-BioID sample revealed 'cell junctions', 'actin binding', and 'cytoskeleton organization' as top terms. Known foot process components were identified and we further uncovered two novel proteins: the tricellular junctional protein Ildr2 and the CDC42 and N-WASP interactor Fnbp11. We confirmed Ildr2 and Fnbp11 are expressed by podocytes and partially colocalize with podocin. Finally, we investigated how this proteome changes with age and uncovered a significant increase in Ildr2. This was confirmed by immunofluorescence on human kidney samples and suggests altered junctional composition may preserve podocyte integrity. Together, these assays have led to new insights into podocyte biology and supports the efficacy of utilizing BioID in vivo to interrogate spatially localized proteomes in health, aging, and disease.

#### 99 Introduction

Kidneys perform vital functions as they filter waste and toxins from the blood and regulate 100 101 body fluid homeostasis. The simplest functional unit of the kidney is the nephron, composed of a 102 blood filter termed the glomerulus connected to a segmented tubule. The glomerulus operates as a 103 'molecular sieve', filtering blood and inhibiting passage of large macromolecules and red blood 104 cells into the nephron tubule. The glomerulus relies on specialized epithelial cells called podocytes. 105 named for their unique cellular morphology with long extruding projections, termed foot processes. 106 Podocyte foot processes wrap around a tuft of fenestrated endothelial capillaries leaving small 107 gaps, or slits, between them. Podocytes undergo morphological changes to their junctional 108 architecture during development to form a specialized barrier between foot processes termed a slit 109 diaphragm.<sup>1-3</sup> The slit diaphragm executes multiple functions including macromolecular filtering in 110 collaboration with the underlying glomerular basement membrane, connection to the actin 111 cytoskeleton to maintain foot process architecture, and signaling that regulates podocyte integrity. 112 The slit diaphragm is distinct from other junctional complexes as it integrates unique structural components as well as components of adherens and tight junctions 4-8. Nearly 50 years ago, a 113 114 zipper-like model for the slit diaphragm was proposed, wherein proteins from neighboring foot processes partially cross the intervening intercellular space and overlap, forming the dense 115 116 protein-rich slit diaphragm structure eloquently visualized by electron microscopy (EM).<sup>9,10</sup> More recent block-face scanning electron microscopy has revealed a 'ridge-like prominence' architecture 117 to podocyte foot processes, formed on the basal surface of the primary foot process.<sup>2,11</sup> These 118 119 investigations underscore the continued advancement in our understanding of podocyte structure 120 and function.

121 When podocytes undergo stress or injury above a threshold, they initiate a response that 122 leads to foot process effacement, loss of slit diaphragms, and proteinuria. Loss of podocyte 123 integrity, observed as effacement, is associated with proteinuric kidney disease<sup>12–14</sup>. This pathology has been described in both acquired and hereditary forms of glomerular disorders or 124 125 podocytopathies.<sup>15</sup> Podocytopathies are a class of kidney diseases in which direct or indirect 126 podocyte injury drives proteinuria or nephrotic syndrome and can ultimately lead to end-stage renal 127 disease (ESRD). Genetic studies have previously identified mutations in numerous podocyte foot 128 process components such as Membrane associated guanylate kinase WW and PDZ Domain 129 Containing 2 (MAGI2)<sup>16</sup>, CD2-associated protein (CD2AP)<sup>17,18</sup>, nephrin (NPHS1)<sup>6,19</sup>, and podocin 130 (NPHS2)<sup>7,20</sup> as causal for nephrotic disease. Additionally, diseases such as diabetes and 131 autoimmune disorders can lead to podocyte injury<sup>21</sup>. While the downstream result is effacement 132 and loss of slit diaphragms, we know little about the temporal changes occurring specifically within 133 the foot process and locally at the slit diaphragm.

134 The identification of slit diaphragm protein complexes with immunoprecipitation followed by 135 mass spectrometry (MS) has uncovered important localized interactions<sup>22,23</sup>. However, the 136 efficiency of these immunoprecipitations is often hindered by harsh conditions required to extract membrane proteins, which can eliminate weaker binding interactions, or by antibodies that may 137 138 disrupt interactions. Additionally, transient interactions may be missed in such experiments. 139 Podocin is known to localize to the slit diaphragm and interacts with both nephrin, Neph1, and 140 Cd2ap.<sup>24–27</sup> Podocin is a member of the stomatin family, containing a central hinge region that 141 integrates into the membrane of the foot process with cytoplasmic N and C termini.<sup>28</sup> Previous 142 studies demonstrated podocin's role in the development of the multiprotein-lipid super complex of the slit diaphragm.<sup>27</sup> Podocin's ability to oligomerize and act as a protein scaffold at slit diaphragms. 143 144 ideally positions it for use as a bait protein in proteomic studies.

145 The discovery and engineering of a promiscuous prokaryotic biotin ligase by Roux and 146 colleagues and concomitant blossoming of –omics technologies in the last decade have laid the 147 groundwork to uncover spatially localized proteomes.<sup>29,30</sup> Proximity-dependent biotin identification, 148 or BioID, utilizes a mutated prokaryotic biotin ligase fused to a bait protein of interest to covalently 149 attach biotin to proteins within the vicinity of a bait protein. The radius of biotinylation can range 150 from ~10nm to 25nm, dependent on the size of the linker between the bait and the biotin ligase. 151 allowing for the biotinylation of both direct and indirect interactors.<sup>29,30</sup> The BioID system requires 152 exogenous addition of biotin for the ligase to covalently biotinylate a proximally located protein. 153 giving the BioID system spatiotemporal control of protein tagging. Additionally, the biotin labels are 154 stable and can withstand harsh isolation conditions, allowing the capture of transient interactors 155 and membrane proteins, respectively. The BioID system has provided plentiful in vitro reports from 156 cell culture models that highlight the power of the system and its ability to discover novel 157 components of even well-documented cellular machinery such as the centrosome and cilium.<sup>31,32</sup> 158 There have been limited in vivo reports of BioID, but it has been used successfully in vertebrates 159 such as zebrafish and mice to identify endogenous interactomes.<sup>32–34</sup> To the best of our knowledge, this approach has not been employed in the mammalian kidney. 160

Here, we utilized gene editing to introduce a smaller, more efficient promiscuous biotin ligase (BioID2) into the endogenous murine *Nphs2* locus to create a fusion protein, hereafter referred to as podocin-BioID (*Nphs2<sup>BioID2</sup>*). Our podocin-BioID model offers the capacity to uncover proteins that localize to the region of the podocyte foot process within the vicinity of podocin in an *in vivo* mammalian system. We were able to identify novel podocyte foot process proteins and furthermore how this proteome changes with age, highlighting the utility of our model for uncovering new interactors as well as disease-associated changes.

#### 169 **Results**

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#### 171 Generation of the *Nphs2<sup>BiolD2</sup>* knock-in mouse model

172 To generate our mouse model, we utilized CRISPR/Cas9 gene editing in combination with homology-directed repair to knock-in the HA-tagged, mutated A. aeolicus biotin ligase (BioID2) in 173 174 frame at the *Nphs2* locus<sup>30</sup>. A single guide RNA (sgRNA) was used to target the stop codon within 175 the eighth exon of Nphs2 to create the fusion. A 13x Glycine/Serine(G/S) flexible linker region was 176 included between Nphs2 and the ligase to provide up to a 25nm reach (Figure 1A)<sup>30</sup>. C57BL/6J 177 zygotes were injected and after screening the resulting animals for a single knock-in with the 178 correct sequence, several founders were identified. A single male founder was utilized for subsequent breeding and expansion of the Nphs2<sup>BiolD2</sup> line. Genotyping E18.5-P0 pups from 179 180 incrosses of *Nphs2<sup>BiolD2/+</sup>* animals identified wildtype, heterozygous, and homozygous offspring at 181 approximately anticipated Mendelian ratios of 29%, 52%, and 19%, respectively (n=73, Figure 1B,C). However, we were typically unable to recover homozygous Nphs2<sup>BiolD2/BiolD2</sup> pups after 1-182 week of birth. Overall, the gross morphology of homozygous pups and kidneys appeared normal at 183 184 P0 and we were unable to determine the specific cause of their death. Due to the early 185 homozygous death, all subsequent experiments for proteomic profiling were performed on heterozygous Nphs2<sup>BioID2/+</sup> animals. To test the function of the podocin-ID fusion and determine if 186 the ligase was able to biotinylate podocyte proteins, we administered 5 mg/kg biotin for 7 187 consecutive days to 8-10 week old Nphs2<sup>BioID2/+</sup> and wild type control mice, a dosage utilized in 188 previous protocols for *in vivo* BioID experiments.<sup>34</sup> Following 1 week of subcutaneous biotin 189 190 injections, kidneys harvested from *Nphs2<sup>BiolD2/+</sup>* mice displayed a pronounced streptavidin signal 191 within glomeruli and specifically podocytes marked by Wt1-positive nuclei (Figure 1D). We 192 occasionally detected streptavidin-positive signal within the tubules which either represents 193 background staining or uptake of free biotin, as controls also displayed this non-glomerular signal 194 pattern. Finally, probing for the HA tag contained within the podocin-BioID fusion shows that the 195 HA signal overlaps with the streptavidin signal, confirming the fusion protein is being specifically 196 translated in podocytes and that the biotin ligase is functional (Figure 1D).

# Podocin-BiolD kidneys display normal nephron morphology and the fusion localizes to the slit diaphragm

199 We further wanted to confirm that the kidneys of animals expressing the podocin-BioID 200 fusion did not display any significant phenotypic differences, most specifically to the nephron. If the 201 fusion protein was not localizing or functioning properly, the animals may display phenotypes 202 associated with Nphs2 knockout animals such as enlarged glomeruli, vacuolated podocytes, and 203 mesangial expansion.<sup>35</sup> Additionally, dilated tubules may indicate abnormal nephron function. 204 Immunostaining kidney sections of wild type, heterozygous, and homozygous animals at E18.5 205 revealed no qualitative differences in glomerular size or proximal tubule dilation (Figure 2A). This 206 further supports that mislocalization or abnormal function of podocin-ID in podocytes is not the 207 likely cause of death in the homozygous animals. Immunofluorescence staining showed a 208 significant overlap between the HA signal and signal from a podocin-specific antibody, supporting 209 that the fusion protein is localizing properly in heterozygous animals and that it is being expressed 210 at similar levels in the homozygous animals (Figure 2A). Importantly, we did not observe the HA 211 antibody signal anywhere else besides the glomerulus.

212 Additionally, we wanted to confirm that the podocin-BioID protein was localizing to the slit 213 diaphragm where podocin is known to interact with other slit diaphragm proteins and play a 214 functional role.<sup>24–27,35</sup> We utilized immunogold labeling in combination with transmission electron microscopy (TEM) to examine the subcellular localization of the podocin-BioID protein (Figure 2B, 215 B'). Nphs2<sup>BiolD2+</sup> kidney sections were stained with an anti-HA antibody followed by a colloidal gold-216 217 AffiniPure secondary. Punctate gold signals were observed near electron dense regions between 218 foot processes, the location of the slit diaphragm (Figure 2B, B' arrowheads). We observed 219 minimal gold signal outside of this region. To confirm that this is the site of normal podocin 220 localization in our control animals, wildtype littermates were probed with a podocin antibody and 221 also showed localization of punctate gold signal near the electron dense region of the slit 222 diaphragm (Figure 2 C, arrowheads).

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#### 224 An enrichment of biotinylated proteins is detected in lysates from *Nphs2<sup>BiolD2/+</sup>* kidneys

225 To enrich our podocin-BioID protein lysates for glomeruli, we surgically isolated the cortex from each kidney of Nphs2<sup>BioID2/+</sup> animals and wildtype controls at 8-10 weeks. Isolated cortex was 226 227 homogenized and lysed to obtain protein lysates from each animal. Protein lysates were applied to 228 magnetic streptavidin coated beads to isolate the biotinylated proteins and an aliguot was removed and tested to validate the efficacy of the biotin ligase. By Western blot analysis, we identified an 229 increase in the number of biotinvlated proteins in our Nphs2<sup>BiolD2/+</sup> sample versus wildtype along 230 231 the full spectrum of molecular weights (Figure 3A). In contrast, few streptavidin labeled, 232 biotinylated, proteins were visible in controls. The few biotinylated proteins observed in wildtype littermates likely represent the endogenous metabolic CoA carboxylases<sup>36</sup>. Due to podocin's ability 233 234 to oligomerize and the heterozygous nature of the mice, we would expect that endogenous, non-235 tagged podocin would be biotinylated as well as podocin-BioID itself. When we probed the Western blot for podocin, we observed two bands within the Nphs2<sup>BiolD2/+</sup> sample: one band at 236 237 approximately 50 kDa, the predicted endogenous podocin molecular weight without the BioID2 tag 238 (Figure 3C, denoted with an asterisk), and a second larger protein, podocin-BioID (Figure 3C, 239 denoted with arrowhead). No bands for podocin or podocin-BioID were detected in controls, as expected. Additionally, the HA signal on the BioID2 protein was only detected in our Nphs2<sup>BioID2/+</sup> 240 241 protein lysates confirming the purity and specificity of our results (Figure 3B). We went on to test if 242 biotin could cross the placental barrier to be delivered to embryonic pups via injection of the 243 pregnant dam. Pregnant dams were injected each day for one week, from E11.5-E18.5, 244 subcutaneously with 5 mg/kg biotin. Pups were then collected at P0. We analyzed the kidney 245 cortex and observed a strong streptavidin signal within the glomeruli of P0, Nphs2<sup>BiolD2/+</sup> mice (Fig 3D, arrowheads). Contrarily, we did not observe signal in wildtype control littermate mice. 246

Together, these data highlight the efficacy and specificity of our podocin-BioID model which can be utilized across a spectrum of ages.

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#### 250 Identification of the podocyte foot process proteome by mass spectrometry profiling

251 Kidney cortices of three 8 to 10-week-old, sex matched mice were collected as one biological replicate, after biotin administration for 1 week. All MS analyses were run in triplicate *i.e.* 252 9 mice per condition, totaling 18 mice (Nphs2<sup>BiolD2/+</sup> vs control littermates) per MS analysis. To 253 254 representatively capture the podocyte foot process proteome, we combined the results of three 255 separate MS analyses, representing 54 total mice (Figure 4). Significant sex differences were not 256 apparent in our studies (Supplemental Tables II-IV). From our proteomics analysis 11 proteins 257 were found across all three MS analyses to have an averaged  $Log_2$  fold change > 2.5. Many of 258 these top proteins including podocin, Kirrel, Tjp1, Pard3, Magi2, Dnd, and Synpo are documented to localize to the slit diaphragm.<sup>37</sup> Yet others including an Immunoglobulin like domain containing 259 260 receptor 2 (Ildr2) and a Formin binding protein 1 like (Fnbp1I, also known as Toca1) were, until now, unreported components of the podocyte foot process/slit diaphragm. Additionally, 6 proteins 261 262 were identified across two MS analyses as significantly unregulated with an average Log<sub>2</sub> fold change  $\geq$ 1.75. These include documented slit diaphragm components Tjp2 and Cd2ap (Figure 263 264 4A). All top 17 proteins had a stringent  $Log_2$  fold change > 1.75. Tables listing the significant foot process proteins identified from each MS analysis are reported in Supplemental Tables II – IV. To 265 add support to our relative Log<sub>2</sub> fold cut off, we assayed an immunoglobulin superfamily adhesion 266 267 molecule, Jam1/F11r,<sup>38</sup> with an observed Log<sub>2</sub> fold expression change of 0.7 (Supplemental Table 268 II). We identified Jam1 expression in cells of the proximal convoluted tubule, abutting the 269 glomerulus, (Supplemental Figure 1) but not within the glomerulus. Potentially, non-glomerular 270 cortex proteins that are biotinylated are also isolated although these appear minimal in our 271 findings. However, this helped establish a relative fold-cutoff for which we start to identify non-272 podocyte proteins.

273 To surmount a complete list of podocyte foot process proteins we established a Log<sub>2</sub> fold 274 change cut off at 1.2 across all three mass spec analyses, and excluded histone, ribosome, and 275 mitochondrial proteins to arrive at a catalog of 54 proteins (Supplemental Table V). Graphically 276 depicting the compiled proteome in a volcano plot with the total ~1400 proteins identified and 277 cataloging proteins with a Log<sub>2</sub> fold change > 1.75 and p-value < 0.05 in green, proteins found to 278 have a p-value < 0.05 in blue and proteins not found to be significant in grev (Figure 4B. 279 Supplemental Table V). The blue dotted line represents a  $-Log_{10}$  (p-value  $\leq 0.05$ ) (Figure 4B). All 280 significantly identified proteins (p < 0.05) had at least two unique peptides identified via MS 281 analysis. The most highly detected and significant proteins from our proteomic profiling, clustering 282 with documented slit diaphragm components, are found in the right scatter of the volcano plot, depicted with green dots (Figure 4B, Supplemental Table V). 283

284 To interrogate our proteomic findings further, the top 54 podocyte foot process proteins 285 were utilized for *in silico* analyses. We input Supplemental Table V into *Qiagen Ingenuity Pathway* 286 Analysis (IPA) and performed a variant effect analysis to compute a proposed interactome based 287 on published literature (Figure 4C). The representative web of interactions from IPA was color coded based on the number of MS analyses each protein appeared in, one (yellow), two (light 288 289 green), and three (dark green) (Figure 4C). Two novel foot process proteins were identified by IPA, 290 Ildr2 and Fnbp1I, to be involved in the interactome with connections to Afadin and actin, 291 respectively (Figure 4C). We added two dotted magenta lines for Ildr2 and Enbp11 potential 292 interactions with podocin (Figure 4C). Unshaded proteins and dashed lines are predicted 293 interactors/interactions from IPA. Our IPA proposed network includes 26 of our 54 input foot 294 process proteins. The IPA interactome identifies three main nodes, *i.e.* junctions, cytoskeleton, and 295 signaling (Figure 4D). We highlighted the respective proteins that contribute to each node for

296 junctions (cyan), cytoskeleton (magenta), and signaling (orange) (Figure 4D). We next input these 297 54 proteins into the Database for Annotation, Visualization, and Integrated Discovery (DAVID) to 298 identify the gene ontogeny (GO) terms that are most highly enriched across our foot process 299 proteome. DAVID analysis identified cytoskeleton protein binding, cell-cell junctions, and actin 300 filament-based processes as the top molecular function, cellular component, and biological processes GO terms, respectively (Figure 5A). Additionally, DAVID and IPA overlap in their 301 302 representation of junctional signaling and actin binding as top GO terms and canonical pathway 303 respectively. Taken together, these findings align with the major functional role roles of podocin 304 and other foot process/slit diaphragm components.

To further decipher the contributions of our top 54 podocyte foot process proteins to junctions and cell-cell contacts we cataloged the protein domains represented within each protein. We utilized binary counting for the presence and absence of a domain. We did not account for the number of same/similar domain(s), *i.e.* SH2/3, within a protein. We identified 7 top protein domains within the catalog of the podocyte foot process proteome that included SH, PDZ, FERM and Ig-like domains among others (Figure 5B). All of these top protein domains align with the nodes of the podocyte foot process proteome, *i.e.* junctions, cytoskeleton, and signaling.

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#### 313 Ildr2 and Fnbp1I are expressed by podocytes and co-localize with podocin

314 Our proteomic analyses uncovered several proteins which have not been previously 315 described to localized to the podocyte foot process or have podocyte-specific functions. We 316 therefore wanted to validate their expression in podocytes and any colocalization with podocin. We 317 utilized immunofluorescence on kidney sections to visualize the location of three proteins in 318 particular: Myozap, Fnbp1I, and Ildr2. First, a known slit diaphragm and foot process component 319 which was identified in our proteomics, Pard3, was assessed for its colocalization with podocin. 320 Pard3 both exhibited positive staining in the glomerulus and overlapped with the HA signal of 321 podocin-BioID (Figure 6A). Pard3 positive staining is also found in the adjacent tubule cells where 322 it helps maintain epithelial integrity (Figure 5A).<sup>39</sup> Myozap regulates cardiac function through Rho-323 dependent activity and interacts with junctional proteins such as ZO-1<sup>40</sup>. It was identified across 324 two of the three MS analyses (Figure 4A) and therefore we decided to investigate its localization. 325 Utilizing an antibody against a synthetic human MYOZAP peptide,<sup>41</sup> we identified strong signal of Myozap in the kidney endothelium, both in the glomerular capillaries and outside of glomerular 326 327 structures (Supplemental Figure 2). Myozap exhibits some degree of overlap with podocin, however, the majority of Myozap protein detection does not colocalize with podocin, suggestive of 328 329 additional roles outside of podocytes (Supplemental Figure 2).

330 We identified Fnbp1I, a documented junctional and actin organizing protein,<sup>42, 43</sup> across all 331 three MS analyses with a Log<sub>2</sub> fold change > 4 (Figure 4). Fnbp11 has been identified in human and mouse podocytes from single cell RNA-seq analysis and linked to podocyte cytoskeleton 332 333 dynamics in vitro, although it has not been documented to colocalize with the slit diaphragm or foot process-associated proteins.<sup>44,45</sup> We found that Fnbp1I co-localizes with podocin in continuous 334 335 stretches within glomeruli observed as a white signal from the Fnbp11 (magenta) and podocin 336 (green) overlap (Figure 6C). In situ hybridization confirms the glomerular expression of Fnpb11 337 (Figure 7C' arrowhead), similar to *Nphs2* (Figure 7A), in addition to a tubular expression pattern at 338 postnatal day 2 (P2).

Ildr2, a member of the B7 superfamily of immunoglobulins, is found highly enriched in the podocyte foot process proteome across all MS profiles with an average Log<sub>2</sub> fold change of 4.9 (Figure 4). Ildr2 is a member of the angulin family and localizes to tricellular junctions of *in vitro* cultured epithelial cells<sup>46</sup>. In the kidney, Ildr2 exhibited a punctate staining pattern that colocalized with podocin in adult mouse glomeruli, reminiscent of the punctate and restricted pattern of tricellular junction staining observed *in vitro* (Figure 6B, 6D arrowheads)<sup>46</sup>. In the z dimension we observe overlap of Ildr2 (magenta) with podocin (green) as white punctate foci denoted by

346 arrowheads (Figure 6D). We further validated *Ildr2* expression within glomeruli via in situ hybridization. Ildr2 expression is found in glomeruli (Figure 7B' arrowhead) and tubule cells of the 347 348 kidney (Figure 7B' arrow) in P2 mice. We went on to verify Ildr2 localization within early renal 349 vesicles, comma, and S-shaped bodies of developing nephrons at E15.5 (Figure 8B, Supplemental 350 Figure 3). Ildr2 is more membranous and contiguous yet with some punctate detection in early 351 embryonic stages (E15.5) both within podocytes and cells of the developing nephron tubule 352 (Figure 8B). Collectively these studies provide evidence that Ildr2 and Fnbp1I are novel podocyte 353 foot process and potentially slit diaphragm components that likely play roles in helping to maintain 354 podocyte architecture.

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#### 356 Ildr2 protein levels increase in both mouse and human glomeruli

357 To determine whether our podocin-BioID model can detect changes to the localized proteome in aging, we performed proteomics on 108-week-old male Nphs2<sup>BiolD2/+</sup> mice and 358 compared the results to our 8-10 week old Nphs2<sup>BiolD2/+</sup> male mice to assess any changes (Figure 359 8A). We identified significant increases in Tip2 and Pkp4 and the largest change (2.76-fold) in Ildr2 360 (Figure 8A). We were unable to detect any significant decrease in proteins at  $\rho < 0.05$ , although 361 362 Fnbp1I and Tns2 showed a decreasing trend. We observed minimal change in podocin when 363 comparing the two age groups (Figure 8A). To assess the human relevance of our findings, we assaved ILDR2 immunofluorescent staining in young (age ~30 years) and aged (91-year-old) 364 human kidney sections. ILDR2 displayed a similar punctate staining pattern in young human 365 glomeruli. In correlation with our findings from the mouse, we found an increase in ILDR2 staining 366 367 in the aged glomeruli. Interestingly, the expression pattern no longer displays a punctate pattern 368 but rather a more diffuse membranous staining pattern (Figure 8C). We quantified the corrected 369 total glomerular florescence (CTGF) by selecting the glomerulus as a region of interest (gROI) then 370 quantifying total florescence intensity and subtracting out the mean background florescence for the 371 glomerular area, *i.e.* total gROI florescence – (glomerular area x mean background florescence) = 372 CTGF, (Figure 9A). We confirmed a significant (p < 0.007) increase in Ildr2 in aged human 373 glomeruli compared to young human kidney tissue (Figure 9A). 374

#### 375 **Discussion**

376 Podocytes are extraordinary epithelial cells of the kidney that intertwine their foot process 377 extensions to establish a cellular junction, the slit diaphragm, that is distinct from other cellular 378 junctions in the human body. The slit diaphragm was identified more than five decades ago as an 379 electron dense region between two podocyte foot processes, visualized beautifully by EM.47,48 380 Many proteins that compose the slit diaphragm when mutated are associated with nephrotic 381 diseases, including nephrin, podocin, Magi2, and Cd2ap.<sup>49</sup> Loss of podocyte integrity is one of the 382 most common clinical observations in kidney disease. End stage renal disease being a top 10 383 cause of death in the US necessitates the need to identify novel components of the slit diaphragm, 384 and how the slit diaphragm changes with disease, for the development of new therapeutic options 385 and biomarkers of disease severity.

386 We utilized a new *in vivo* biochemical tool to interrogate the proteome of the podocyte foot 387 process via knock-in of a BioID moiety, generating our podocin-BioID model. The sensitivity and 388 advantage of BioID is that it allows for weak and transient interactions to be identified, in addition 389 to withstanding harsh isolation conditions.<sup>6</sup> Further, the biotin-streptavidin bond is one of the 390 strongest known non-covalent interactions, enhancing the isolation of biotinylated proteins via 391 streptavidin-coated magnetic bead. We validated our model recapitulates normal podocin 392 localization by TEM and showing podocin-BioID localizes normally to the slit diaphragm. Activity of 393 the biotin ligase was confirmed through probing kidney sections and lysates with streptavidin and 394 finding an enrichment of biotinylated proteins. Our proteomics profiling identified more than 50

395 candidate molecules with significant enrichment in the podocyte foot process,  $Log_2 \ge 1.2$ . Within 396 this dataset we were able to identify novel candidates not found in previous investigations of the slit diaphragm proteome<sup>22,23</sup>. Additionally, we were able to identify changes to the localized 397 398 proteome that occur with aging. It would be interesting to compare the proteome across additional 399 stages such as during development. Podocytes begin as a columnar epithelium with only tight and 400 adherens junctions which remodel and later mature to form the specialized slit diaphragm 401 complex.<sup>16</sup> With the ability of biotin to cross the placental barrier, the proteome could be compared 402 from development to maturity to identify changes that may help inform how these specialized 403 junctions are formed.

404 The proteomics analysis revealed potential nuances of the podocyte foot process that are 405 still under investigation. We identified 11 proteins that repeated over all three MS profiles. 406 However, there were more than 30 proteins that only surfaced in a single MS analysis and split 407 across the three MS analyses. While this may be due to experimental variability, such as in the 408 isolation of proteins, this may also highlight some differences in biological activity in the foot 409 process. The dynamics, variability, and protein turnover in the podocyte slit diaphragm is relatively 410 unknown, Pointing to the relative variability of one documented component, Synaptopodin (Synpo). 411 across the three MS analyses the respective Log<sub>2</sub> fold changes were observed at 1.0, 2.9, and 4.8 412 which indicates potential protein turn over, variability in the slit diaphragm components, or potential variability from MS to MS. Further still, a second documented slit diaphragm protein, Cd2ap, was 413 only identified in two of the MS profiles. These cofounding issues made a single MS analysis a 414 415 limited view, or snapshot, of the proteins present within the podocyte foot process. By combining 416 three separate MS analysis we have uncovered a more complete profile of the podocyte foot 417 process and slit diaphragm. Many of the top candidates identified in the podocyte foot process 418 proteome are well documented slit diaphragm components including Kirrel, Nphs2, Par3, Magi1/2, 419 Tjp1/2/3, Dnd, Synpo, and Cd2ap. Furthermore, our in silico analysis utilizing DAVID highlights 420 anticipated GO terms, including actin binding, cell-cell junctions, adherens junctions, and 421 cytoskeleton organization that align with podocyte function. We conclude the podocin-BioID model 422 provides a spatial specific approach to identify proteins that are notoriously difficult to isolate.

423 We cannot rule out biotin delivery and metabolism may limit some proteins from being 424 detected. One report depicts a time lapse of biotin administration in vivo with positive streptavidin 425 enrichment for as long as 18-hour after biotin administration.<sup>50,51</sup> However, biotin requires an 426 accessible primary amine within a peptide to biotinylate. Proteins with little open structure or a 427 limited number of lysine residues could be missed by the labeling strategy. The possibility that 428 proteins were missed is brought to light by the inability to identify nephrin (Nphs1) across any of 429 the mass spec analyses run. One rational for the inability to identify nephrin, a known slit 430 diaphragm molecule that has been shown to interact with podocin, posits that its C-terminal 431 domain is only minimally assessable to podocin, while the remaining majority of the 180 kDa 432 protein is spanning the extracellular space of the slit diaphragm.<sup>27</sup> Therefore, while there are significant advantages to the BioID system over traditional immunoprecipitations followed by MS, it 433 434 is also subject to missing important interactions.

435 One of the novel foot process candidates we identified, Fnbp1I, has multiple roles in 436 microtubule binding, cell polarity, motility, actin organization, junctional localization and 437 signaling.<sup>42,43</sup> Based on these roles, we hypothesize that Fnbp11 similarly helps maintain the slit 438 diaphragm and foot process integrity through associations with the actin cytoskeleton and links to 439 the slit diaphragm. Single cell RNA-seg analyses have previously found *Endp11* expression in 440 human<sup>44</sup> and mouse podocytes and tubule cells.<sup>45</sup> From the Kidney Interactive Transcriptomics 441 (KIT), *Fnbp1* is most highly expressed in podocytes compared to the tubule cells of the human kidney.<sup>52</sup> Further, *Fnbp11* is also expressed in the zebrafish pronephros<sup>53</sup> with a similar pattern to 442 443 other zebrafish tight junction proteins such as ZO-1/2<sup>54</sup>, however the specific expression of *Fnbp11* 

within the zebrafish podocytes remains unknown. With the conservation of *Fnbpl1* expression in
 the renal system across vertebrates, it likely plays an important role in helping maintain epithelial
 integrity in the various cells of the nephron, including podocytes.

447 Ildr2 was identified in the podocyte foot process/slit diaphragm proteome across all three 448 MS profiles with a Log<sub>2</sub> fold change > 4.5 and a p-value < 0.007. Ildr2 belongs to the B7/CD28 449 family of proteins, encompassed by the Ig superfamily (IgSF), with pivotal roles in self-tolerance.55 450 immunomodulation and maintenance of peripheral lldr2 evinces immunomodulatory activity, wherein administration of Ildr2 as a fusion protein (ILDR2-Fc) 451 452 rebalances immune homeostasis, which leads to an amelioration of autoimmune disease states in 453 mouse models of rheumatoid arthritis, type I diabetes, and relapsing-remitting multiple sclerosis.<sup>55-</sup> 454 56,57 Recently, an Ildr2 blocking antibody, BAY 1905254, was generated to block the 455 immunosuppressive activity of Ildr2 for cancer immunotherapy.<sup>57</sup> BAY 1905254 promotes T-cell activation in vitro and enhances antigen-specific T-cell proliferation and cytotoxicity in vivo, and is 456 457 currently in phase I clinical trials.<sup>57</sup> Ildr2 presents an unexplored niche in podocyte biology centered around the function of podocytes in immune cell modulation. Yet, the ability for immune cells such 458 459 as T-cells to interreact with podocytes is challenged by the podocyte cellular environment and the 460 GBM. Therefore, whether Ildr2 could play an immune related role in podocytes remains tentative.

461 Ildr2/ILDR2 is found to be specifically expressed in mouse and human podocytes from publicly available databases including KIT.<sup>44,58</sup> *Ildr2* was also identified as a Wt1 transcriptional 462 target in podocytes by ChIP-seq.<sup>59</sup> In early embryonic and postnatal mouse kidneys, *Ildr2/*Ildr2 is 463 expressed and localized to tubule cells and podocytes. However, in the adult kidney Ildr2 464 465 localization becomes restricted to podocytes. At the timepoints of podocyte development, we 466 observe a more membranous and contiguous pattern of Ildr2 in comma and s-shaped bodies, with 467 some foci of heightened detection. In mature glomeruli, Ildr2 is restricted to punctate foci that 468 colocalize with podocin. Ildr2 has been identified at the site of tricellular tight junctions (tTJ) in 469 murine retinal pigment epithelium and *in vitro* within EpH4, a mouse mammary gland cell line.<sup>46,56</sup> 470 Additionally. Ildr2 has recently been identified to interact with Afadin in human embryonic kidney 471 cells (HEK293).<sup>60</sup> Afadin is also reported to localize to the site of specialized tricellular junctions and mediate mechanotransduction.<sup>61</sup> The role of mechanical strain present from the underlying 472 473 fluid flow sheer stress on these specialized tTis and recruitment of specific proteins to these 474 domains remains unresolved. Many proteins identified in the podocyte foot process proteome are 475 documented junctional proteins containing classified SH, PDZ, FERM, transmembrane, EF-Hand, 476 and Ig-like domains. Yet, Ig-like domain containing proteins were a considerably smaller 477 population. It is intriguing to speculate that Ig-like domains or a combination of PDZ, SH, and Ig-478 like domain aids in the recruitment of specific proteins to specialized junctions such as tTJs and the 479 slit diaphragm. Further still, the requirement of these proteins for managing stress such as from 480 changes in fluid flow from the fenestrated endothelium and conversely the potential requirement of 481 mechanostress for recruitment of proteins to these tTJs and slit diaphragms remains an area in 482 need of further investigation.

483 In our aged podocin-BioID mice as well as our aged human kidney sample, we identified an 484 increase in IIdr2 within podocytes. The pattern of IIdr2 is more membranous with some punctate 485 foci in embryonic renal structures yet becomes detectable only as punctate foci as the mouse 486 matures. Proteomic profiling of 108-week-old, aged mice identify a significant increase in Ildr2 487 detection. We validated Ildr2 significantly increases in aged (91 yo) human glomeruli via guantifying the corrected total glomerular florescence of young and aged human glomeruli. We 488 489 further noted that in the aged human glomeruli Ildr2 is detected higher in zones where podocin is 490 less detected and conversely where podocin is detected high Ildr2 is less abundant. Aged human 491 glomeruli present an Ildr2 pattern that more closely resembles the mouse embryonic stage with a 492 contiguous pattern rather than the punctate pattern of younger mice. We hypothesize that Ildr2 is a

tight junction component and increasing Ildr2 protein in aged podocytes helps maintain podocyte
 integrity with age. The additional increase of tight junction and desmosome associated proteins
 Tjp2 and Pkp4, respectively, in our aged mouse proteome further supports this hypothesis.

In this study, we developed an innovative tool for the identification of novel proteins within spatially restricted podocyte foot process/ slit diaphragm. Current efforts are now aimed at uncovering proteins differentially regulated during slit diaphragm development, aging, and in renal disease. It is crucial we uncover the changing composition of the podocyte slit diaphragm and foot process as it holds potential for new therapeutic treatment targets to either preserve or prevent the loss of podocyte integrity in kidney disease.

# 502503 Methods

504 Genetic CRISPR/Cas9 engineering of the Nphs2 locus to append a BioID2 moiety

505 CRISPR/Cas9 targeting, and donor vector design were performed by UNC Chapel Hill Animal 506 Models Core. *Benchling* software was used to identify Cas9 short guide RNAs (sgRNA) overlapping the podocin (Nphs2) stop codon. Guide RNAs were cloned into a T7 promoter vector 507 508 followed by *in vitro* transcription and spin column purification. Functional testing was performed by 509 transfecting a mouse embryonic fibroblast cell line with sqRNA and Cas9 protein (produced and 510 purified by the UNC Protein Expression Core). The sqRNA target region was amplified from 511 transfected cells and analyzed by T7 endo1 nuclease assay (New England Biolabs). sgRNA. Nphs2-g78B (protospacer sequence 5'-gCCATTCGCCTATAACAT-3'; lower case g indicates 512 513 heterologous quanine added at 5' end of native sequence for efficient T7 in vitro transcription) was 514 selected for genome editing in embryos. Nphs2 was amplified from adult mouse kidney cDNA and 515 cloned into the MCS-BioID2-HA plasmid (plasmid was a gift from Kyle Roux; Addgene plasmid # 74224 ; http://n2t.net/addgene:74224 ; RRID:Addgene\_74224).30 A donor vector was subsequently 516 517 constructed from the Nphs2-BioID2-HA plasmid with the following features: (1) a 1018 bp 5' homology arm encompassing sequence immediately 5' of the Nphs2 stop codon including 276 bp 518 519 coding sequence from *Nphs2* exon 8. (2) a 267 bp in-frame Glycine/Serine-rich linker sequence. 520 (3) a 696 bp coding sequence for the humanized biotin ligase of *A. aeolicus* with a R40G mutation in the catalytic domain (BioID2)<sup>30</sup>, (4) HA Tag, (5) 2X Stop codon, (6) a FRT site, and (7) a 1019 bp 521 522 3' homology arm, beginning at the Nphs2-sgRNA cut site. The donor vector was designed to 523 produce a final knock-in allele which would produce a fusion protein of podocin C-terminally linked 524 to BioID2.

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#### 526 Embryo microinjection

527 C57BL/6J zygotes were microinjected with 400nM Cas9 protein, 25 ng/µl sgRNA and 20 ng/µl 528 supercoiled double-stranded donor plasmid (Mix1), or 200 nM Cas9 protein, 12.5 ng/µl sgRNA and 529 10 ng/ul donor plasmid (Mix2). Injected embryos were implanted in pseudopregnant B6D2F1 recipient females. Fourteen resulting pups (9 from Mix1 and 7 from Mix2) were screened by PCR 530 531 for the presence of the knock-in allele. One female (Founder #2) and one male (Founder #6) were 532 positive for the correct single-copy knock-in allele. Founders #2 and #6 were mated to wild-type 533 C57BL/6J animals for transmission of the knock-in allele. Both founders transmitted the correct 534 knock-in allele to offspring. Injections, genotyping of founders, and off-site targeting analysis was 535 performed by UNC Chapel Hill Animal Models Core.

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#### 537 Genotyping

A mouse tail or ear clip was taken and dissociated with Viagen DirectPCR Lysis Reagent (Mouse Tail) containing 10  $\mu$ g/mL proteinase K incubated at 55°C overnight and denatured at 95 °C for 10 min. PCR was run with (T<sub>annealing</sub> = 63.5°C), elongation for 40 sec, for 35 cycles. Primer sequences utilized for genotyping are Common Forward: 5'-CTTTTGTCCTCCCGGCAA-3',

542 podocin WT Reverse 5'-TGCATGTAGCCATCTTGTGACT-3', *Nphs2<sup>BiolD2</sup>* Reverse 5'-543 CTGCCCTTGGTCTGTCTGTC-3'.

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563

#### 545 Kidney cortex isolation and lysis

546 Mice were raised, housed, and handled in accordance with IACUC protocol number 19-183.0/22-136.0. 8–10-week-old mice were injected subcutaneously with 5 mg/kg biotin every day 547 548 for one week. We surgically isolated the cortex of the kidney to enrich for the glomerular fraction, 549 using a scalpel in cold sterile phosphate buffered saline (PBS). A single biological sample was 550 composed of the isolated kidney cortex of three same sex mice for a total of six kidney cortexes 551 per sample (2 kidneys per animal x 3 mice). The sample was then homogenized via a glass 552 Dounce homogenizer. Samples were centrifuged at 4 °C, 5000 × g for 10 minutes. Supernatant 553 was decanted and the tissue pellets snap frozen in liquid nitrogen and stored at -80 °C. Care was 554 taken to utilize sterile Eppendorf tubes that had not been autoclaved as we identified a 555 polyethylene glycol contaminant in a preliminary MS analysis that may arise from autoclaving 556 plastic utilized. Samples were removed from the freezer and allowed to equilibrate on ice for 2 557 hours. Lysis buffer (8M urea. 50mM Tris-HCl pH 7.4. 500mM NaCl. 2.5mM EDTA. 2.5mM EGTA. 558 1.5mM MgCl2, 1.5mM DTT, 0.25% NP-40, 1% SDS, 1x protease inhibitors (Roche cOmplete Mini 559 EDTA-free, Sigma)) were then added to resuspend the pellet and allowed to nutate at 4°C for one hour. The protein homogenate was sonicated (3 pulses for 10 sec at 30% duty) and centrifuged 560 (4°C, 12000 rpm for 10 min). The supernatant was then removed for subsequent quantification and 561 562 analysis.

564 Biotinylated protein capture

565 Kidney cortex-isolated protein lysates were serial diluted and run-in triplicate on a 96-well 566 plate reader at 590nm wavelength (Synergy HT, BioTek). A standard curve of bovine serum 567 albumin (BSA) was utilized as a control. Protein concentrations of each lysate were calculated, and 10mg of crude protein lysate was loaded with 100µl of streptavidin-coated magnetic beads 568 569 (Dynabeads MyOne Streptavidin C1, Invitrogen). Samples containing beads and protein lysate 570 were rotated end over end at 4°C overnight. Supernatant was removed using a magnetic strip, and 571 the bead-captured fraction was washed once with lysis buffer. After the first wash with lysis buffer 572 containing 1% SDS and 0.25 % NP-40 diminishing amounts of detergents were utilized in 573 subsequent washes until there was no detergents remaining in the wash buffer (after 4 washes). 574 Beads were then washed 3 times in ABC solution (50mM ammonium bicarbonate, 8.0 pH) and 575 sent to UNC Chapel Hill Hooker Proteomics Core. All procedures were performed the same for our 576 aged murine cohort with the exception that 5mg of crude protein lysate was loaded onto beads.

After the last wash buffer step, 50µl of 50mM ammonium bicarbonate (pH 8) containing 1 µg trypsin (Promega) was added to beads overnight at 37°C with shaking. The next day, 500ng of trypsin was added then incubated for an additional 3h at 37°C with shaking. Supernatants from pelleted beads were transferred, then beads were washed twice with 50ul LC/MS grade water. These rinses were combined with original supernatant, then acidified to 2% formic acid. Peptides were desalted with peptide desalting spin columns (Thermo Scientific) and dried via vacuum centrifugation. Peptide samples were stored at -80°C until further analysis.

- 584
- 585 LC/MS/MS analysis

Each sample was analyzed by LC-MS/MS using an Easy nLC 1200 coupled to a QExactive HF (Thermo Scientific). Samples were injected onto an Easy Spray PepMap C18 column (75µm id × 25cm, 2µm particle size) (Thermo Scientific) and separated over a 120 min method. The gradient for separation consisted of a step gradient from 5 to 36 to 48% mobile phase B at a 250 nl/min flow rate, where mobile phase A was 0.1% formic acid in water and mobile phase B consisted of 0.1% formic acid in ACN. The QExactive HF was operated in data-dependent mode where the 15 most

intense precursors were selected for subsequent HCD fragmentation. Resolution for the precursor scan (m/z 350–1700) was set to 60,000 with a target value of  $3 \times 10^6$  ions, 100ms inject time. MS/MS scans resolution was set to 15,000 with a target value of  $1 \times 10^5$  ions, 75ms inject time. The normalized collision energy was set to 27% for HCD, with an isolation window of 1.6m/z. Peptide match was set to preferred, and precursors with unknown charge or a charge state of 1 and  $\geq 8$  were excluded.

#### 599 Western blot

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600 Biotinylated proteins attached to streptavidin coated bead were eluted off with excess biotin 601 [200 mM] in 200 mM Tris HCl pH: 6.8, 40% glycerol, 8 % beta mercaptoethanol, 2 % SDS, 0.04 % 602 bromophenol blue at 95°C for 30 min). Bead slurry mix was placed on a magnet and isolated 603 biotinylated lysate was collected for subsequent analysis. Protein sample fractions were removed 604 and loaded into a Novex WedgeWell gel (4-20% Tris-Glycine gradient gel, Invitrogen, XP04202), 605 and run in 25mM Tris-HCl, 190mM Glycine, 0.1% SDS, pH 8.3. Proteins were transferred from gel 506 to nitrocellulose membrane in 25mM Tris-HCI, 190mM Glycine, and 20% methanol. The membrane 507 was blocked in 3% bovine serum albumin in 1x TBST (Tris base Saline Solution (25mM Tris-HCI pH 7.5, 150mM NaCl) with 0.1% Tween-20, for 1 hour at room temperature. Primary antibodies 608 509 (Rabbit anti-HA tag (Cell Signaling, 3724S [1:500], mouse anti-NPHS2 (Proteintech, 20384-1-AP) 610 [1:500], Rabbit anti-podocin (Invitrogen, PA5-79757) [1:500], Streptavidin-HRP (Cell Signaling, 3999S) [1:1000] were applied in 3% BSA+ TBST. Membrane plus primary antibodies were allowed 611 512 to incubate overnight at 4°C with gentle nutation. Following 3x10 min washes with TBST, 613 membranes were probed with secondary antibodies conjugated to horseradish peroxidase (HRP) 614 (donkey and rabbit -HRP [1:500] or Goat anti mouse-HRP [1:500] and incubated for one hour at 615 room temperature Membranes were developed with enhanced chemiluminescence substrate 516 (ECL) and visualized on an iBright FL1000 (Invitrogen).

#### 618 Proteomics data analysis

619 Raw data were processed using the MaxQuant software suite (version 1.6.12.0) for identification and label-free quantitation.<sup>62</sup> Data were searched against an Uniprot Reviewed 620 Mouse database (downloaded January 2021, containing 17,051 sequences) using the integrated 521 622 Andromeda search engine. A maximum of two missed tryptic cleavages was allowed. The variable 623 modification specified was oxidation of methionine, N-terminal acetylation, and phosphorylation of 624 Ser/Thr/Tyr. Label-free quantitation (LFQ) was enabled. Results were filtered to 1% FDR at the 625 unique peptide level and grouped into proteins within MaxQuant. Match between runs was 626 enabled. Data were filtered in Perseus, then imported into Argonaut for normalization, imputation 627 and statistical analysis.63

We combined three separate MS analyses and averaged the Log<sub>2</sub> fold changes and respective *p*values of the top 17 proteins identified across all three MS profiles. QIAGEN Ingenuity Pathway Analysis (IPA) was utilized to identify putative interactomes.

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#### 632 Immunofluorescence

633 Kidneys were harvested in cold filter sterilized PBS and fixed at 4°C for 1 hr in 4% 634 paraformaldehyde (PFA) in PBS. Samples were washed twice in 1x PBS, placed in 30% sucrose 635 overnight, and subsequently embedded in OCT. Kidneys were cut in 12µm sections on a Leica 636 Cryostat CM 1850. Tissue sections were blocked in 3% donkey serum, 1% bovine serum albumin (BSA), 1x PBS, 0.1 % TritonX-100 for 45 min at room temperature. Slides were incubated in 637 638 primary antibodies (see antibodies utilized below, Supplemental Table I) diluted in blocking buffer 639 for 1-2 hr. Slides were then rinsed 3x5 min in 1x PBST, after which they were incubated with Alexa 640 Fluor labeled secondary antibodies (Invitrogen) diluted (1:1000) in blocking buffer for 1 hr at room

temp. The slides were then rinsed 2 x 5 min with 1x PBST, 1x3-5 min with 1x PBS + 1ng/mL DAPI and mounted in ProLong Gold antifade reagent (Invitrogen).

643 For double labeling tissue with two antibodies both raised in rabbit we utilized a Zenon 644 double labeling kit (Z25302, Invitrogen) with rabbit anti-Fnbp1I, rabbit anti-Ildr2, rabbit anti-podocin, 645 and rabbit anti-HA antibodies. The weaker of the two primary rabbit antibodies, typically Ildr2 and 646 Fnbp11, was diluted in block solution then incubated for 90 min at room temperature. Slides were washed 3-6x with 1xPBS +0.25% Triton-X 100 and subsequently incubated for 1hr at room 647 temperature with the appropriate Alexa Fluor-488 labeled secondary antibody diluted 1:1000 in 648 649 blocking buffer. The second primary rabbit antibody was mixed at a 1:2.5 ratio with the Zenon 650 fluorophore-647. Note the two fluorophores utilized are on opposite ends of the florescence 651 spectrum. The slides were then fixed in 4% PFA for 5 min at room temperature. And all 652 subsequent steps were carried out per normal immunofluorescence procedures as above. Images 653 were acquired utilizing a Zeiss 880 confocal microscope equipped with Airyscan super-resolution 654 and spectral imaging on Zen Microscopy Suite version 2.3 Sp1 that is part of the UNC Hooker 655 656 Imaging Core. Z-stack images were acquired in 1µm steps for Ildr2 and Fnbp11.

#### 657 Immunogold electron microscopy

Mouse kidneys were fixed for 1 hour at room temperature in 4% PFA in 0.15M sodium phosphate 658 buffer pH 7.4 (PB) or 2% PFA + 0.5% glutaraldehyde in 0.15M PB and immediately processed for 659 660 LR White resin embedding. Samples were washed in 0.15M PB, 3x10 minutes and dehydrated using a Pelco BioWave Pro Microwave (Ted Pella, Inc.) as follows: at 40°C (750 watts): 30% 561 662 ethanol in water (ETOH) -1 min, 50% ETOH in water-1 min, 75% ETOH in water-1 min, and at 663 40°C (450 watts): 100% ETOH-1 min, 100% ETOH-1 min, 100% ETOH-1 min. Microwave infiltration and embedment were carried out using the following schedule: 1 part 100% ETOH:2 664 665 part LR White resin-10 min at 40°C (350 watts), 2 exchanges of 100% LR White resin for 10 min at 50°C (350 watts). Samples were transferred to 00 gelatin capsules filled with fresh LR White resin 566 667 and polymerized overnight at 55°C; the temperature was adjusted to 60°C for 6 hours and allowed to complete polymerization at room temperature for 72 hrs. Blocks were trimmed to the tissue and 668 669 1.0µm sections were cut using Leica Ultracut UCT (Leica Microsystems, Inc.) and a Diatome 670 diamond knife (Electron Microscopy Sciences). Sections were mounted on glass slides, stained 671 with 1% toluidine blue O in 1% sodium borate, and regions with glomeruli were selected and 672 trimmed.<sup>64</sup> Ultrathin sections (90 nm) were cut and mounted on formvar / carbon-coated 200 mesh 673 nickel grids. Before immunostaining, the sections were hydrated by floating the grids section-side 674 down on drops of deionized water. Sections were blocked with Aurion Goat Blocking Solution for 675 15 min and transferred to  $15\mu$  drops of the primary antisera diluted at [1: 500] for anti-HA and [1: 676 250] for anti-Podocin in 0.05M TBS+0.2% BSA-Ac, pH 7.6 (Rabbit anti-HA, Cell Signaling, 37245; rabbit anti-Podocin, Invitrogen, PA5-79757). Sections were incubated overnight at 4°C followed by 677 678 4x10 min washes in TBS/BSA-Ac to remove unbound antibody. The grids were incubated in a 679 12nm Colloidal gold-AffiniPure goat anti-rabbit IgG (H+L) secondary antibody (Jackson Immuno, 680 Lot #148041) diluted at 1:50 in TBS/BSA-Ac for 2 hours at room temperature.<sup>65</sup> After 3 washes in 681 TBS/BSA-Ac and 3 washes in 0.15M PB, grids were post-fixed for 10 min in 1% glutaraldehyde in 682 0.15M PB followed by 3 washes in deionized water. The grids were stained with 4% aqueous uranyl acetate for 5 min for additional contrast. Samples were observed using a JEOL JEM-1230 683 684 transmission electron microscope operating at 80kV (JEOL USA INC.), and images were taken using a Gatan Orius SC1000 CCD camera with Gatan Microscopy Suite version 3.10.1002.0 685 686 software (Gatan, Inc.). Sample prep and imaging was performed at the UNC Microscopy Services 687 Laboratory, Department of Pathology and Laboratory Medicine.

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#### 591 In situ hybridization

Wildtype C57/BI6J kidneys at postnatal day 2 (P2) were dissected in cold molecular grade 692 593 PBS (RNAse/DNAse free) and fixed in 4% PFA for 30 min at room temperature on a rocking 694 platform. Kidneys were washed in molecular grade PBS and placed in 30% sucrose. Kidneys were **695** embedded in OCT and 20µm sections were cut on a cryostat (Leica CM1850). Tissue sections **596** were fixed in 4% PFA and washed with 1x PBS. The tissue was permeabilized with proteinase K at 10µg/mL for 15 min and subsequently washed with 1x PBS. Tissue was then washed in an 697 acetylation solution (0.1 % HCl, 0.375 % Acetic Anhydride, and 0.75 % Triethanolamine in H<sub>2</sub>0) for **598** <u> 599</u> 10 min with stirring. Tissue sections were then subsequently washed 3x3min with PBS, rinsed 1x5min in 0.85 % NaCl, 1x5min in 70 % ethanol, and 1x5min in 95 % ethanol prior to riboprobe / 700 701 hybridization application. Antisense Digoxygenin (DIG) labeled riboprobes were hybridized to the 702 tissue in a solution containing 50% deionized formamide, 2 sodium citrate pH 4.5, 1% SDS, 50µg/mL heparin, 50µg/mL yeast tRNA overnight in a humidity chamber at 68°C. The following 703 704 day, specimens were treated with successive washes of sodium citrate buffer as described 705 previously,<sup>66</sup> and blocked in 10% heat inactivated sheep serum (HISS), with 2% Roche blocking reagent (BR) in malic acid buffered solution with 0.1% tween-20 (MABT). Anti-DIG-Alkaline 706 707 Phosphatase (AP) antibody was applied in 1% HISS, 2% BR in MABT overnight at 4°C. The 708 following day the slides were washed in MABT and 100mM NaCl, 100 mM Tris-HCl pH 9.5, 50mM 709 MgCl<sub>2</sub>, and 0.1 % Tween-20 (NTMT) with 2mM Levamisole to inhibit endogenous alkaline 710 phosphatase activity. Digoxigenin-UTP (Roche 53119620) labeled riboprobes were amplified from 711 cDNA libraries collected from mouse kidney tissue with the following primers:

712 Podocin forward primer: TGACGTTCCCTTTTTCCATC, Podocin reverse primer with T7 713 underlined: CAGTGAATTG<u>TAATACGACTCACTATAGGG</u>CTGTGGACAGCGACTGAAGA.

714Ildr2forward:GGAGAATCCTTGGGCandIldr2reverse:715CAGTGAATTGTAATACGACTCACTATAGGGGGTACCCGGCCTTGGCwerepreviously716published<sup>66</sup>.Fnbp1Iforwardprimer:GCTGAATGACAATTGTGTGAACandFnbp1Ireverse:717CAGTGAATTGTAATACGACTCACTATAGGGGCTGTGCAAGTCCAAGTGTCTTC.

718

#### 719 Human kidney tissue

Human kidney tissue samples were de-identified (young) or directly donated (91-year-old) and did not necessitate IRB approval. Young, normal kidney tissue was obtained from the UNC Tissue Procurement Facility as a frozen block. A small piece of tissue was removed with a razorblade, fixed for 10 minutes in 4% paraformaldehyde and subsequent processing for cryosectioning and immunofluorescence carried out as for the mouse (described above). For the old kidney tissue, a fresh sample was excised from the kidney cortex and fixed for 10 minutes in 4% paraformaldehyde prior to processing similar to the young tissue for immunofluorescence.

#### 728 Abbreviations

4.1 protein, Ezrin, Radixin and Moesin (FERM), Corrected Total Glomerular Florecence (CTGF), E
and F helix–Hand (EF-Hand), End Stage Renal Disease (ESRD), Kidney Interactive
Transcriptomics (KIT), Gene Ontogeny (GO), Glomerular Basement Membrane (GBM), Mass
Spectrometry (MS), PSD-95, Disc large, and ZO – 1 (PDZ), Src homology (SH),

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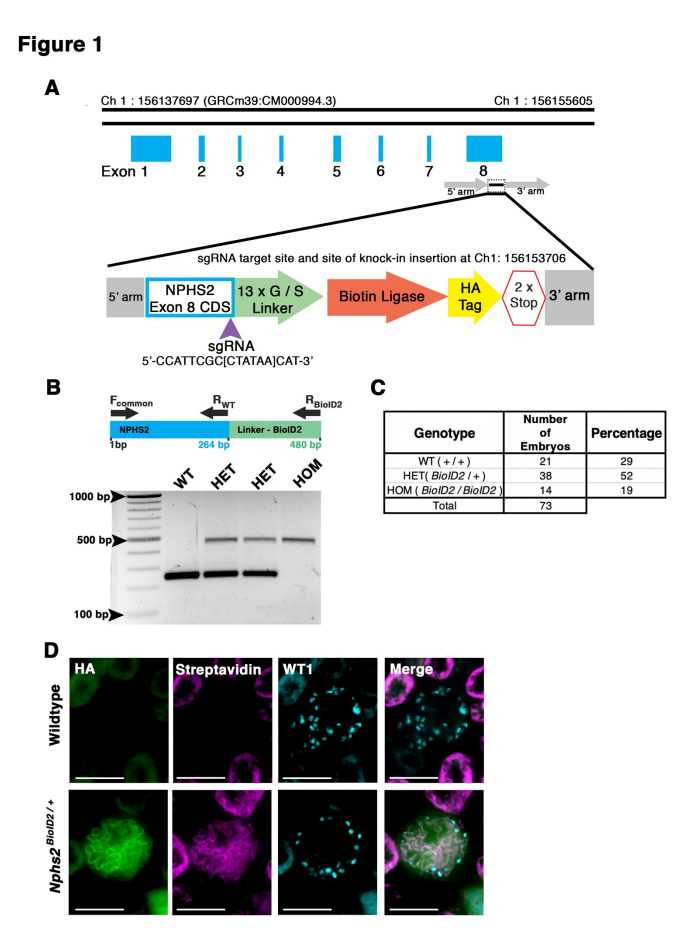
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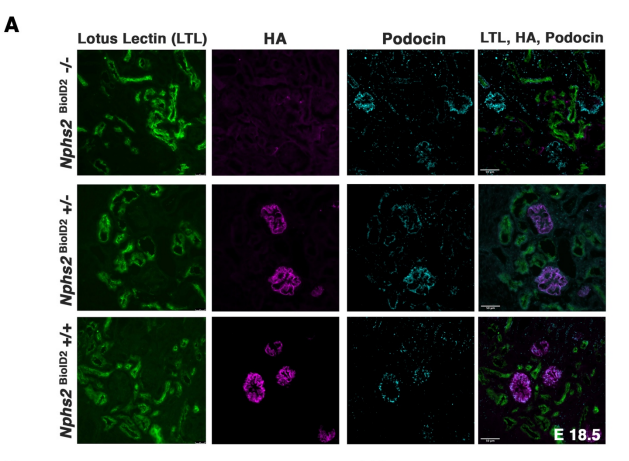
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972 Figure 1. Generation of a knock-in *Nphs2<sup>BiolD2</sup>* mouse line via CRISPR/Cas9 genome editing of the Nphs2 locus. (A) Schematic of the CRISPR/Cas9 genome editing strategy utilized to 973 generate the *Nphs2<sup>BiolD2</sup>* mouse line. A small guide RNA (sgRNA) targeting the stop codon in exon 974 8 of *Nphs2* (purple arrowhead) was combined with a donor vector containing the knock-in cassette 975 976 (zoom view) to induce homologous recombination and integrate the BioID2 moiety containing a 13x Glycine/Serine (G/S) linker, biotin ligase, and HA tag into the Nphs2 locus. (B) Genotyping 977 978 strategy (top panel) to identify genotype of mice as wildtype (single band at 264 base pairs (bp)), 979 heterozygous (two bands; one at wildtype size of 264 bp and a second that amplifies the BioID2 980 linker region giving a band at 480 bp), or homozygous (single band at the 480 bp). (C) Genotyping 981 of 73 embryos at E18.5-P0, verify an approximate Mendelian ratio of genotypes being recovered 982 (25:50:25). (D) IF analysis of 8–10-week-old adult mice injected with biotin illustrate an enrichment of streptavidin detected within the glomerulus of Nphs2<sup>BiolD2</sup> mice and absence of streptavidin 983 signal in control age matched C57BL/6J littermate mice. The HA signal from the BioID2 moiety 984 985 closely overlaps with streptavidin, observed as white overlap in the merged image. Wilms Tumor (WT1) is utilized as a podocyte marker to delineate the glomerular boundaries. Scale bar: 50 µm 986 987

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### Figure 2.



В

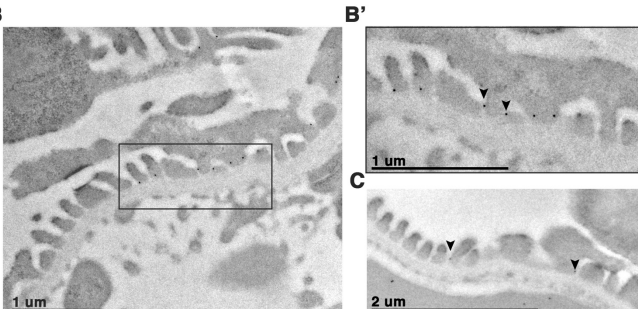
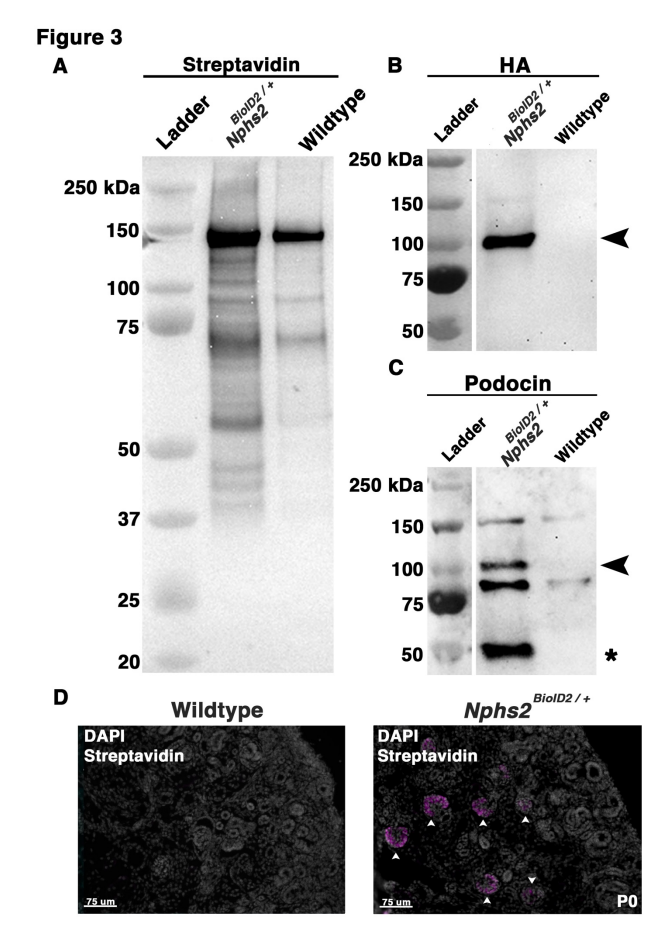


Figure 2. Nphs2<sup>BiolD2</sup> animals present normal kidney morphology with podocin-BiolD localized to podocytes and specifically the slit diaphragm. (A) E18.5 littermates of wildtype, heterozygous (Nphs2<sup>BiolD2/+</sup>), and homozygous (Nphs2<sup>BiolD2/BiolD2</sup>) animals were analyzed for gross kidney morphology and localization of podocin-BioID. The proximal tubule marker lotus lectin (LTL, green) show normal tubule architecture without observation of nephron tubule dilation in heterozygous (Nphs2<sup>BiolD /+</sup>), and homozygous (Nphs2<sup>BiolD2 / BiolD2</sup>) animals. The HA tag (magenta) which is part of the BioID2 moiety was observed to tightly overlap with podocin (cyan) specifically in the glomerulus. The relative size and number of glomeruli was unchanged between control, heterozygous, and homozygous animals. Further, the HA signal was not identified outside of the glomeruli. Scale bar 50 µm (B) Nphs2<sup>BiolD2/+</sup> localizes to the podocyte slit diaphragm. Utilizing transmission electron microscopy (TEM) and immunogold labeling of 4-week-old murine kidney samples, we detect the anti-HA signal as punctate dots strung just above the GBM adjacent to the podocyte foot process within electron dense regions where the slit diaphragm bridges neighboring foot processes. (B') Magnified view of boxed region from panel (B) depicting podocyte foot processes and the electron dense regions between them where the HA signal localizes as dark spherical dots from immunogold labeling, denoted with arrowheads. Scale bar 1 μm (C) Wildtype littermates were immunogold labeled for podocin and similarly display localization of podocin at electron dense regions between podocyte foot processes, highlighted with arrowheads. Scale bar 2 um.



042 Figure 3. Biotin administered Nphs2<sup>BiolD2/+</sup> mice present an enrichment of biotinylated proteins specifically within glomeruli. (A) A significant enrichment of streptavidin labeled, 043 044 biotinvlated, proteins are observed across the entire molecular weight spectrum in Nphs2<sup>BioID2/+</sup> 045 protein lysates compared to wildtype control littermates, injected with biotin. Streptavidin bead 046 purified lysates were subjected to protein separation and probed for streptavidin. Wildtype age matched, 8–10-week-old, littermate controls present few streptavidin conjugated bands, 047 presumably endogenous metabolic carboxylases. However, *Nphs2<sup>BiolD2/+</sup>* lysates exhibit numerous 048 specific biotinvlated protein bands across the entire molecular weight spectrum. (B) Nphs2<sup>BiolD/+</sup> 049 050 lysates blotted with an anti-HA antibody present robust and specific detection of the HA signal only within the Nphs2<sup>BiolD2/+</sup> sample, indicative of lysate purity. The HA signal is evident at approximately 051 100 kDa (arrowhead) in Nphs2<sup>BiolD2/+</sup> lysates. (C) Nphs2<sup>BiolD2/+</sup> samples probed for podocin show 052 053 two specific bands for both BioID2-tagged podocin and endogenous podocin. Adult 8-10-week-old Nphs2<sup>BiolD2/+</sup> mice blotted for podocin present a 50kDa band for the endogenous unmodified 054 055 podocin protein, marked with an asterisk (\*) and a second larger band at approximately 100 kDa for the HA tagged podocin-BioID of *Nphs2*<sup>BioID2/+</sup> mice (arrowhead). (D) P0 pups collected from 056 057 pregnant dams injected with 5 mg/kg of biotin every day from E11.5 to E18.5, and without injection of newborn pups, manifest strong streptavidin detection within glomeruli. Immunofluorescence 058 059 analysis reveals an enrichment of streptavidin signal (magenta) specifically within the glomeruli (arrowheads) of *Nphs2<sup>BiolD2/+</sup>* kidneys compared to wildtype. Scale bar 75 µm. 060

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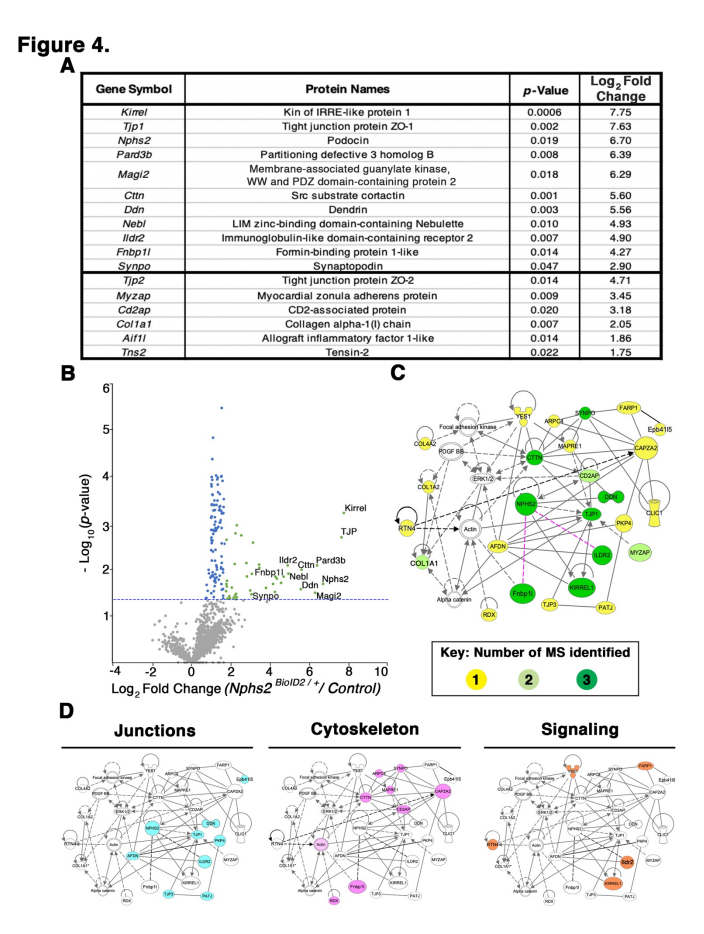
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077 Figure 4. Proteomics profiling of the podocyte foot process identifies documented slit diaphragm components and novel candidates with unexplored podocyte function. (A) Three 078 079 separate MS analyses were combined, and the average Log<sub>2</sub> fold change and respective p-values 080 were averaged to produce a list of the top foot process proteins. The top 11 proteins were 081 identified across all three MS profiles. Six additional proteins were identified across two of the three MS analyses, denoted following the thick black bar. Podocin was a top protein identified and 082 083 known to oligomerize, indicating our MS proteomic profiling was effective. Additionally, many of the 084 top proteins uncovered, *i.e.*, Kirrel, Dnd, Tjp1/2, Magi2, Pard3, Synpo, and Cd2ap are documented 085 podocyte slit diaphragm components. (B) A volcano plot depicting approximately 1400 proteins 086 identified across all three MS analyses. Green dots represent the 40 proteins identified across 087 three separate MS profiles as having a Log<sub>2</sub> fold change  $\geq$  1.7, and *p*-value  $\leq$  0.05, blue dots 088 denote all proteins identified to have a significant p-value < 0.05, and grey dots are proteins with a 089 *p*-value > 0.05. The top 11 proteins consistently uncovered across all MS analyses are embedded 090 with gene symbols in the plot. The blue dotted line represents a p-value < 0.05. (C) Network 091 topology was generated utilizing Qiagen Ingenuity Pathway Analysis (IPA) on the top 54 proteins 092 from our cumulative proteomic profiles. IPA produces a proposed web of relationships from 093 published literature with most of our top proteins represented, (11 of 17), with documented 094 connections to other foot process and slit diaphragm components. The color intensities of each protein (vellow to green) represent the number of MS analyses from which each protein was 095 096 identified. Darkest green shade=all three MS analyses, lighter green shade=two of three MS 097 analyses, and yellow=a single MS analysis. Dotted lines and unshaded proteins represent 098 predicted interactions / interactors from IPA analysis, respectively. Magenta lines are novel 099 proteins identified in this study to be present within the podocyte foot process. (D) Dissecting the 100 IPA network, identifies three central nodes representing junctions, cytoskeleton, and signaling. 101 Each respective node is color coded with junctions in cyan, cytoskeleton in magenta, and signaling in orange. These computational network topologies and nodes tightly align with podocyte structure 102 103 and function.

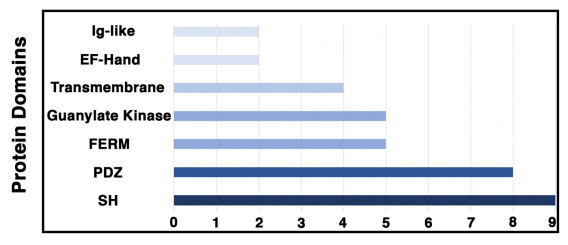
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## Figure 5.

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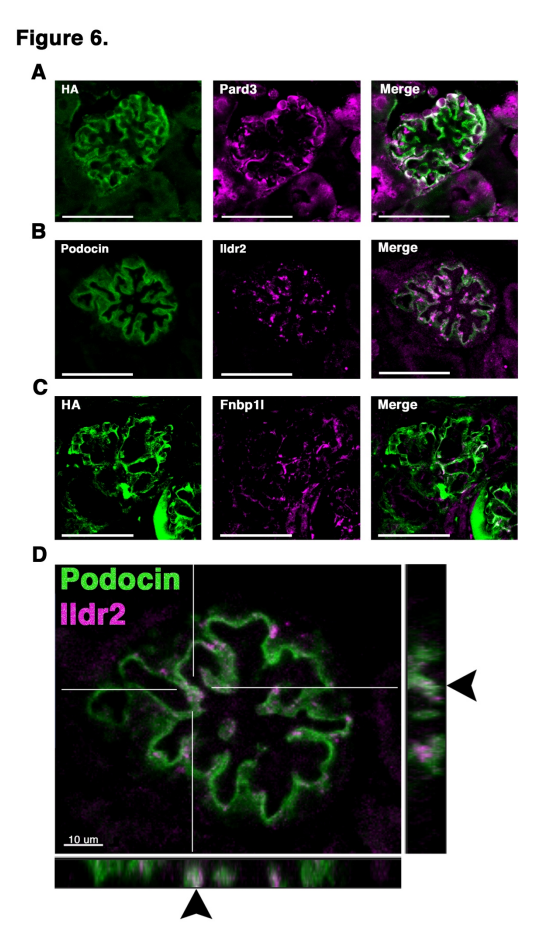
Gene Ontology (GO) Category	Cluster	Enrichment Score	Counts	p-Value
Molecular Function (MF)	Cytoskeletal Protein Binding	4.53	12	2.20 x 10
Molecular Function (MIT)	Actin Binding	4.00	6	7.80 x 10
Cellular Component (CC)	Cell-Cell Junction	4.42	12	7.70 x 10
Cellular Component (CC)	Adherens Junction		10	7.90 x 10
Biological Process (BP)	Cytoskeleton Organization	2.61	14	3.20 x 10
biological Flocess (DP)	Actin Filament-Based Process	2.01	12	8.70 x 10

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Number of Proteins w/ Domain from Foot Process Proteome

106 Figure 5. Analysis of the podocyte foot process proteome identifies the cytoskeleton, cellcell junctions, and actin-based processes as the top GO categories with respective protein 107 108 domains that align with these functions. The Database for Annotation. Visualization, and 109 Integrative Discovery (DAVID) was utilized for GO clustering analysis of the top 54 proteins 110 identified to have a Log<sub>2</sub> fold change > 1.20, and p-value < 0.05 (Supplemental Table V). (A) Overlap in GO readout was observed across the three separate GO categories analyzed. 111 112 molecular function, cellular component, and biological process, with cytoskeleton, actin, and 113 cellular junctions being the top hits. The respective top clusters within each GO category are listed. 114 alongside the relative enrichment score, number of proteins, and *p*-values. (B) Protein domain analysis of the top 54 podocyte foot process proteins provides evidence for protein-protein 115 116 interactions and likely scaffold and protein-protein complex formation. Each protein's respective 117 protein domains were binary counted for presence or absence within the proteomics profile. The top seven protein domains identified from the podocyte foot process proteome consist of a Src 118 119 homology (SH) (n = 9 proteins), PSD-95, Disc large, and ZO – 1 (PDZ) (n = 8), 4.1 protein, Ezrin, Radixin and Moesin (FERM) (n = 5), Guanylate Kinase (n = 5), Transmembrane (n = 4), E and F 120 121 helix - Hand (EF - Hand) (n = 2), and an Immunoglobulin - like (lg - like) domain (n = 2). 122 Cumulatively, these 7 domains represent ~ 50 % of the top proteins identified from the foot process 123 proteome (26 / 54).



125 Figure 6. Two novel foot process candidates, Ildr2 and Fnbp1I, localize to podocytes and overlap with podocin in the foot process. IF analysis of three top candidates identified from the 126 proteomics profiling, Pard3, Ildr2, and Fnbp1I, were detected within glomeruli. (A) Pard3 127 (magenta), a known component of the podocyte foot process, co-localizes with HA signal of 128 129 podocin-BioID (green) producing a white signal in overlap. (B) Ildr2 (magenta) displays a punctate localization pattern within the glomerulus and overlaps (white) with podocin (green). (C) Fnbp11 130 (magenta) co-localizes (white) with HA signal of podocin-BioID (green) in podocytes. Scale bars in 131 A-C: 50 µm (D) A merged image of podocin and Ildr2 through a confocal z-projection identifies 132 punctate localization and overlap (white, arrowheads) between podocin (green) and Ildr2 133 (magenta) visualized in the z-plane in the left and bottom panels. Scale bar in D 10  $\mu$ m. 134

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# Figure 7. A' A 南小田 Nphs2 B' В lldr2 C' С Fnbp1l P

Figure 7. In situ hybridization confirms glomerular and some tubule cell expression of Ildr2 137 and *Fnbp11* in P2 kidneys. P2 kidney sections were hybridized with antisense riboprobes against 138 Nphs2 (podocin), Ildr2, and Fnbp11. (A) Nphs2 (purple) displays strong expression specifically in 139 glomeruli at P2. The rectangular box in (A) is enlarged in (A') to highlight expression of Nphs2 only 140 141 in glomeruli, arrowheads denote example glomeruli (B) Ildr2 transcripts (purple) are identified within glomeruli and tubules. The rectangular box in (B) is enlarged in (B') to denote expression of 142 143 *Ildr2* in glomeruli, arrowhead, and in some tubules, black arrow. (C) *Fnbp11* (purple) is identified in 144 both glomeruli and tubules. The block rectangular box in (C) is enlarged in (C') to denote expression of *Fnbp11* in glomeruli, arrowhead, and tubules, arrow. Scale bars in A-C: 500 μm. 145 Scale bars in A'-C': 100 µm. 146

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## Figure 8.

Gene Symbol	Protein Names	BiolD (Young) vs WT Log <sub>2</sub> Fold Change	BioID (Aged) vs WT Log <sub>2</sub> Fold Change	<i>p</i> -Value				
lldr2	Immunoglobulin-like domain-containing receptor 2	2.76	3.53	0.001				
Tjp2	Tight junction protein ZO-2	2.71	3.31	0.01				
Pkp4	Plakophilin-4	2.27	2.94	0.01				
Nphs2	Podocin	2.42	2.36	0.10				
Fnbp1l	Formin-binding protein 1-like	3.22	2.12	0.10				
Tns2	Tensin-2	1.30	0.36	0.10				

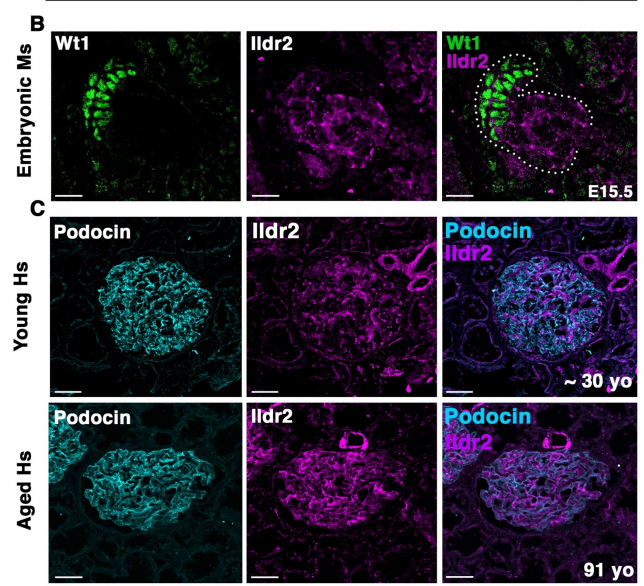
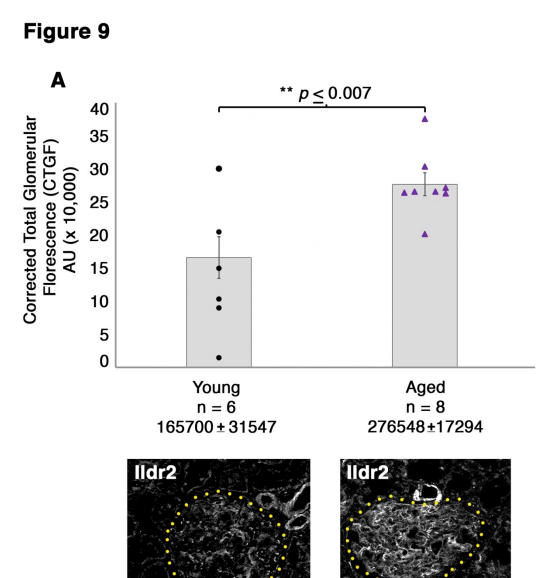


Figure 8. Ildr2 is detected in developing nephron structures and presents increased 154 detection in aged mouse and human glomeruli compared to young. (A) Proteomic profiling of 155 two aged (108 week old) male *Nphs2<sup>BiolD2/+</sup>* mice identifies a significant increase in Ildr2. Tip2, and 156 Pkp4 protein levels compared to 8-10 week old Nphs2<sup>BiolD2/+</sup> mice. Table lists 6 proteins with 157 158 relative detection in "young" 8-10 week old Nphs2<sup>BiolD2/+</sup> male mice vs "aged" 108 week old *Nphs2<sup>BiolD2/+</sup>* male mice. All protiens were detected with at least 2 razor unique peptides with their 159 respective *p*-values in far right column. (B) Ildr2 is expressed in early developing nephron 160 structures including the renal vesicle, comma, and s-shaped bodies with podocytes denoted by the 161 162 Wt1-positive nuclei. Ildr2 (magenta) is observed as membranous along the comma / s-shapped 163 body with some punctate foci in E15.5 mouse kidney cortex, outlined with white dots. Scale bar: 20 164 μm. (C) Young and aged (91 yo) human kidney tissue was probed of Ildr2 via IF. Young human 165 glomeruli display punctate foci detection of Ildr2 (magenta) similar to 8-10 week old mice. 166 However, aged (91 yo) male glomeruli evince an increase in Ildr2 detection within glomeruli. Furthermore, where Ildr2 looks to be more highly detected in these aged glomeruli podocin 167 168 appears less detectable. Scale bar: 50 µm.

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181 Figure 9. Aged 91 yo glomeruli exhibit a significant increase in corrected total glomerular florecence (CTGF) compared to young human glomeruli. We observe a significant (p < 0.007) 182 increase, of approximatly 67%, in CTFG in aged human Ildr2 detection compared to young human 183 tissue. CTGF was calculated by selecting the gomeruli as the ROI then measuring the total 184 185 florescence within the area and subtracting out the sum of the area by the average background florecence, *i.e.* Total gROI florecence – (Area of glomerulus x mean of background florescence) = 186 187 CTGF. Two example glomeruli are displayed and outlined in yellow dots to depict the respective ROI utilized to calculate CTGF. Scale bar: 50 µm. 188

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# Supplemental Table I.

Antidody	Dilution [IF]	Dilution [WB]	Company
Goat anti - Collagen IV	[1:500]		Sigma-Aldrich, AB769, Lot 2930544
Mouse (IgG1) anti - Myozap	supernatant (straight)		Progen, clone 517.67, 651169
Rabbit anti - F11r (Jam1)	[1:50]		ABclonal, A1241, Lot 0005180101
Rabbit anti - Fnbp1I (Toca-1)	[1:100]		Bethyl Laboratories, A303-469A, Lot 1
Rabbit anti - HA	[1:500]	[1:1000]	Cell Signaling, clone C29F4, 3724S
Rabbit anti - Ildr2	[1:250]		BiCell Scientific Inc, 00303
Rabbit anti - Ildr2	[1:50]		Lifespan BioSciences LS-C205504
Rabbit anti - Ildr2	[1:50]		Invitrogen, PA5-46481, Lot WE327570
Rabbit anti - Pard3	[1:250]		Sigma-Aldrich, 07-330, Lot 2615671
Rabbit anti - podocin	[1:500]	[1:1000]	Invitrogen, PA5-79757, Lot WL3458408
Rabbit anti - WT1	[1:500]		Abcam, ab89901, LotGR3365362-7
Lotus Lectin	[1:1000]		Vector Laboratories, FL-1321, Lot ZC0914
Streptavidin - APC	[1:1000]		BD Pharmingen, 554067 Lot 7040924
Streptavidin - HRP		[1:5000]	Cell Signaling, 3999S, Lot 8

- Supplemental Table I. List of antibodies with respective company identifiers and 199 concentrations for immunoflorecence (IF) and western blot (WB) analyses utilized in this 200
- 201 investigation.

# Supplemental Table II.

Gene Symbol	Majority Protein IDs	Protein Names	Razor + Unique Peptides	Log <sub>2</sub> Fold Change	p - Value
Nphs2	Q91X05	Podocin	12	10.63	0.00003
Tjp1	P39447	Tight junction protein ZO-1	85	7.40	0.001
Magi2	Q9WVQ1	Membrane-associated guanylate kinase, WW and PDZ domain-containing protein 2	33	7.37	0.001
Kirrel	Q80W68	Kin of IRRE-like protein 1	20	7.06	0.001
Pard3b	Q9CSB4	Partitioning defective 3 homolog B	32	5.34	0.00002
Ddn	Q80TS7	Dendrin	12	5.33	0.00003
lldr2	B5TVM2	Immunoglobulin-like domain-containing receptor 2	11	4.51	0.009
Cttn	Q60598	Src substrate cortactin	15	4.01	0.0003
Nebl	Q9DC07	LIM zinc-binding domain-containing Nebulette	11	3.65	0.003
Tip2	Q9Z0U1	Tight junction protein ZO-2	31	3.63	0.021
S100g	P97816	Protein S100-G	4	3.06	0.034
Fnbp1l	Q8K012	Formin-binding protein 1-like	7	2.92	0.010
Capza2	P47754	F-actin-capping protein subunit alpha-2	8	2.82	0.014
Atp6v1e1	P50518	V-type proton ATPase subunit E 1	9	2.30	0.039
Canx	P35564	Calnexin	9	2.26	0.042
Mvzap	Q3UIJ9	Mvocardial zonula adherens protein	7	2.09	0.0001
Kpnb1	P70168	Importin subunit beta-1	2	1.96	0.041
Dars	Q922B2	AspartatetRNA ligase, cytoplasmic	7	1.88	0.029
Cndp2	Q9D1A2	Cytosolic non-specific dipeptidase	6	1.83	0.041
Pls3	Q99K51	Plastin-3	5	1.76	0.029
Sult1d1	Q3UZZ6	Sulfotransferase 1 family member D1	5	1.73	0.038
Uqcr10	Q8R111	Cytochrome b-c1 complex subunit 9	3	1.67	0.048
Uba52	P62984; P62983; P0CG49; P0CG50	Ubiguitin-60S ribosomal protein L40	5	1.60	0.041
Sec23a	Q01405	Protein transport protein Sec23A	7	1.59	0.030
Cisd2	Q9CQB5	CDGSH iron-sulfur domain-containing protein 2	2	1.46	0.026
Naca	Q60817:P70670	Nascent polypeptide-associated complex subunit alpha, muscle-specific form	2	1.46	0.049
Ca15	Q99N23	Carbonic anhydrase 15	3	1.44	0.050
Qars	Q8BML9	Glutaminyl-TRNA Synthetase 1	2	1.40	0.039
Rpl35	Q6ZWV7	60S ribosomal protein L35	4	1.37	0.038
Agk	Q9ESW4	Acylglycerol kinase, mitochondrial	5	1.37	0.039
Scp2	P32020	Non-specific lipid-transfer protein	13	1.36	0.039
Aimp1	P31230	Endothelial monocyte-activating polypeptide 2	2	1.26	0.026
Dhrs7b	Q99J47	Dehydrogenase/reductase SDR family member 7B	5	1.20	0.040
Fabp4	P04117	Fatty acid-binding protein, adipocyte	2	1.17	0.024
Mpv17	P19258	Protein Mpv17	2	1.17	0.012
Sf3b1	Q99NB9	Splicing factor 3B subunit 1	4	1.10	0.039
Dkc1	Q9ESX5	H/ACA ribonucleoprotein complex subunit 4	4	1.06	0.003
Por	P37040	NADPH-cytochrome P450 reductase	11	1.05	0.003
	Q8CC35		6	1.02	0.031
Synpo Acox1	Q9R0H0	Synaptopodin Peroxisomal acyl-coenzyme A oxidase 1	24	1.02	0.054
Cisd 1	Q91WS0	CDGSH iron-sulfur domain-containing protein 1	6	0.99	0.015

Supplemental Table II. Proteomic profiling of cohort 1, 8-10 week old *Nphs2*<sup>BiolD2</sup> male mice, identifies 41 significantly detected podocyte foot process proteins. MS analysis of biotin administered male Nphs2<sup>BioID2/+</sup> mice identifies Podocin, Tip, Magi, Kierrel, and Pard3 as top 5 proteins respectively. Table list 41 significant proteins ( $p \le 0.05$ ) at a Log<sub>2</sub> cut off 1.0 with their respective *p*-values in the far-right column. Podocin was identified as the top hit and is documented to oligomerize with itself adding confidence that the MS analysis was successful. All proteins were identified with at least 2 razor unique peptide sequences *i.e.*, two peptide sequences aligned only to that protein.

# Supplemental Table III.

Gene Symbol	Majority Protein IDs	Protein Names	Razor + Unique Peptides	Log₂ Fold Change	p - Valu
Kirrel	Q80W68	Kin of IRRE-like protein 1+D3:D62	15	10.19	0.001
Tjp1	P39447	Tight junction protein ZO-1	85	9.88	0.003
Pard3b	Q9CSB4	Partitioning defective 3 homolog B	26	9.62	0.016
Ddn	Q80TS7	Dendrin	11	8.99	0.005
Magi2	Q9WVQ1	Membrane-associated guanylate kinase, WW and PDZ domain-containing protein 2	24	8.61	0.011
Nebl	Q9DC07	LIM zinc-binding domain-containing Nebulette	7	7.75	0.021
lldr2	B5TVM2	Immunoglobulin-like domain-containing receptor 2	9	7.42	0.010
Cttn	Q60598	Src substrate cortactin	14	7.08	0.020
Nphs2	Q91X05	Podocin	10	7.05	0.023
Fnbp1l	Q8K012	Formin-binding protein 1-like	5	6.65	0.008
Magi1	Q6RHR9	Membrane-associated guanylate kinase, WW and PDZ domain-containing protein 1	6	5.89	0.002
Tjp2	Q9Z0U1	Tight junction protein ZO-2	30	5.78	0.006
Myzap	Q3UIJ9	Myocardial zonula adherens protein	4	4.80	0.018
Synpo	Q8CC35	Synaptopodin	4	4.79	0.043
Khsrp	Q3U0V1	Far upstream element-binding protein 2	5	4.69	0.014
Mapre 1	Q61166	Microtubule-associated protein RP/EB family member 1	4	4.56	0.019
Yes1	Q04736	Tyrosine-protein kinase Yes	8	4.34	0.015
Tmem65	Q4VAE3	Transmembrane protein 65	4	4.25	0.030
Cd2ap	Q9JLQ0	CD2-associated protein	5	4.15	0.019
Ap1g1	P22892	AP-1 complex subunit gamma-1	6	3.85	0.018
Ewsr1	Q61545	RNA-binding protein EWS	2	3.83	0.038
Oxsm	Q9D404	3-oxoacyl-[acyl-carrier-protein] synthase, mitochondrial	4	3.77	0.033
Erlin2	Q8BFZ9	Erlin-2	6	3.44	0.008
Epb4115	Q8BGS1	Band 4.1-like protein 5	9	3.13	0.003
Hsd3b4	Q61767	3 beta-hydroxysteroid dehydrogenase type 4	9	2.99	0.028
Dhtkd1	A2ATU0	Probable 2-oxoglutarate dehydrogenase E1 component DHKTD1, mitochondrial	7	2.93	0.017
Taco1	Q8K0Z7	Translational activator of cytochrome c oxidase 1	2	2.64	0.045
Aif1I	Q9EQX4	Allograft inflammatory factor 1-like	3	2.31	0.001
Gstt2	Q61133	Glutathione S-transferase theta-2	4	2.24	0.002
Sorbs1	Q62417	Sorbin and SH3 domain-containing protein 1	18	2.21	0.036
Tns2	Q8CGB6	Tensin-2	12	2.20	0.007
Farp1	F8VPU2	FERM, RhoGEF and pleckstrin domain-containing protein 1	7	2.13	0.040
Tjp3	Q9QXY1	Tight junction protein ZO-3	7	2.12	0.043
SIc25a42	Q8R0Y8	Mitochondrial coenzyme A transporter SLC25A42	9	1.98	0.021
Mybbp1a	Q7TPV4	Myb-binding protein 1A	13	1.92	0.037
Acnat1	A2AKK5	Acyl-coenzyme A amino acid N-acyltransferase 1	4	1.81	0.046
Epb41/1	Q9Z2H5	Band 4.1-like protein 1	8	1.79	0.023
Col1a1	P11087	Collagen alpha-1(I) chain	8	1.73	0.002
Col4a2	P08122	Collagen alpha-2(IV) chain;Canstatin	5	1.69	0.006
Nomo1	Q6GQT9	Nodal modulator 1	17	1.68	0.014
Col1a2	Q01149	Collagen alpha-2(I) chain	10	1.65	0.001
Dao	P18894	D-amino-acid oxidase	11	1.63	0.002
Arpc4	P59999	Actin-related protein 2/3 complex subunit 4	2	1.62	0.007
Ugcrh	P99028	Cytochrome b-c1 complex subunit 6, mitochondrial	2	1.61	0.019
Atp5l	Q9CPQ8	ATP synthase subunit g, mitochondrial	4	1.59	0.002
Patj	Q63ZW7	Pals1-associated tight junction protein / InaD-like protein	6	1.53	0.003
Dhrs 1	Q99L04	Dehydrogenase/reductase SDR family member 1	8	1.53	0.0003
Apmap	Q9D7N9	Adipocyte plasma membrane-associated protein	14	1.52	0.003
Lad1	P57016	Ladinin-1	14	1.49	0.012
Rtn4	Q99P72	Reticulon-4	8	1.49	0.012

Supplemental Table III. Proteomic profiling of cohort 2, 8-10 week old Nphs2<sup>BiolD2</sup>male mice, identifies 50 significantly detected podocyte foot process proteins. MS analysis of Nphs2<sup>BiolD2/+</sup> biotin administered mice identifies Kirrel, Tjp, Pard3, Ddn, and Magi as the top 5 proteins detected respectively, with Podocin detected in the top 10. Table list 50 significant proteins (p < 0.05) at a Log<sub>2</sub> cut off > 1.49 with respective p-values in the far-right column. One additional immunoglobulin domain containing protein, Junction adhesion molecule 1 (Jam1 / F11r) was also identified within this MS analysis with a Log<sub>2</sub> fold change of 0.7, p < 0.05. All proteins were detected with at least two razor unique peptide sequences to the specific protein identified.

- 27.

# Supplemental Table IV.

Gene Symbol	Majority Protein IDs	Protein Names	Razor + Unique Peptides	Log <sub>2</sub> Fold Change	p - Value
Kirrel	Q80W68	Kin of IRRE-like protein 1	18	5.99	0.027
Cttn	Q60598	Src substrate cortactin	14	5.70	0.034
Tjp1	P39447	Tight junction protein ZO-1	90	5.62	0.050
Pard3b	Q9CSB4	Partitioning defective 3 homolog B	38	4.21	0.050
Nebl	Q9DC07	LIM zinc-binding domain-containing Nebulette	7	3.41	0.051
Fnbp1l	Q8K012	Formin-binding protein 1-like	7	3.22	0.025
Magi2	Q9WVQ1	Membrane-associated guanylate kinase, WW and PDZ domain-containing protein 2	30	2.89	0.044
Synpo	Q8CC35	Synaptopodin	10	2.88	0.044
lldr2	B5TVM2	Immunoglobulin-like domain-containing receptor 2	11	2.76	0.003
Tjp2	Q9Z0U1	Tight junction protein ZO-2	21	2.71	0.037
Col1a2	Q01149	Collagen alpha-2(I) chain	3	2.68	0.046
Nphs2	Q91X05	Podocin	13	2.42	0.035
Col1a1	P11087	Collagen alpha-1 (I) chain	5	2.37	0.011
Ddn	Q80TS7	Dendrin	13	2.35	0.021
Pkp4	Q68FH0	Plakophilin-4	13	2.27	0.033
Cd2ap	Q9JLQ0	CD2-associated protein	3	2.20	0.034
Rdx; Msn	P26043; P26041	Radixin; Moesin	2	1.74	0.015
Afdn	Q9QZQ1	Afadin	2	1.70	0.030
Hspe1	Q64433	10 kDa heat shock protein, mitochondrial	4	1.66	0.036
Aif11	Q9EQX4	Allograft inflammatory factor 1-like	5	1.42	0.027
Hist1h4a	P62806	Histone H4	9	1.36	0.014
Tns2	Q8CGB6	Tensin-2	3	1.30	0.038
Khk	P97328	Ketohexokinase	2	1.25	0.027
Clic1	Q9Z1Q5	Chloride intracellular channel protein 1	2	1.21	0.027
Hist2h2bb	Q8CGP2;Q8CGP1	Histone H2B type 1	6	1.04	0.049
Acadl	P51174	Long-chain specific acvI-CoA dehvdrogenase, mitochondrial	7	1.01	0.025

Supplemental Table IV. Proteomic profiling of cohort 3, 8-10 week old *Nphs2<sup>BiolD2</sup>* female mice, identifies 28 significantly detected podocyte foot process proteins. MS analysis of biotin administered female *Nphs2<sup>BiolD2/+</sup>* mice identifies Kirrel, Cttn, Tjp, Pard3b, and Nebl as the top 5 proteins detected. Table list 28 significant proteins ( $p \le 0.05$ ) at a Log<sub>2</sub> cut off 1.0 with respective *p*-values in the far-right column. Many documented podocytes SD proteins are identified, *i.e.* Podocin, giving indication the podocin-BiolD model is successful. All proteins were detected with at least two razor unique peptide sequences to the specific protein identified.

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# Supplemental Table V.

Gene Symbol	Protein Names	<i>p</i> -Value	Log <sub>2</sub> Fold Change	
Kirrel	Kin of IRRE-like protein 1	0.0006	7.75	
Tjp1	Tight junction protein ZO-1	0.002	7.63	
Nphs2	Podocin	0.019	6.70	
Pard3b	Partitioning defective 3 homolog B	0.008	6.39	
Magi2	Membrane Associated Guanylate Kinase, WW And PDZ Domain Containing 2	0.018	6.29	
Cttn	Src substrate cortactin	0.001	5.60	
Ddn	Dendrin	0.003	5.56	
Nebl	LIM zinc-binding domain-containing Nebulette	0.010	4.93	
Ildr2	Immunoglobulin-like domain-containing receptor 2	0.007	4.90	
Fnbp1l	Formin-binding protein 1-like	0.014	4.27	
Synpo	Synaptopodin	0.047	2.90	
	Tight junction protein ZO-2	0.014	4.71	
Tjp2				
Myzap	Myocardial zonula adherens protein	0.009	3.45	
Cd2ap	CD2-associated protein	0.020	3.18	
Col1a1	Collagen alpha-1(I) chain	0.007	2.05	
Aif 1I	Allograft inflammatory factor 1-like	0.014	1.86	
Tns2	Tensin-2	0.022	1.75	
Magi1	Membrane Associated Guanylate Kinase, WW And PDZ Domain Containing 2	0.002	5.89	
Khsrp	Far upstream element-binding protein 2	0.014	4.69	
Mapre1	Microtubule-associated protein RP/EB family member 1	0.019	4.56	
Yes1	Tyrosine-protein kinase Yes	0.015	4.34	
Tmem65	Transmembrane protein 65	0.030	4.25	
Ap1g1	AP-1 complex subunit gamma-1	0.018	3.85	
Erlin2	Erlin-2	0.008	3.44	
		_		
Epb4115 S100g	Band 4.1-like protein 5 Protein S100-G	0.003	3.13	
			3.06	
Capza2	F-actin-capping protein subunit alpha-2	0.014		
Atp6v1e1	V-type proton ATPase subunit E 1	0.039	2.30	
Pkp4	Plakophilin-4	0.033	2.27	
Canx	Calnexin	0.042	2.26	
Sorbs1	Sorbin and SH3 domain-containing protein 1	0.036	2.21	
Farp1	FERM, RhoGEF and pleckstrin domain-containing protein 1	0.040	2.13	
Tjp3	Tight junction protein ZO-3	0.043	2.12	
Mybbp1a	Myb-binding protein 1A	0.037	1.92	
Cndp2	Cytosolic non-specific dipeptidase	0.041	1.83	
Epb41l1	Band 4.1-like protein 1	0.023	1.79	
Pls3	Plastin-3	0.029	1.76	
Rdx	Radixin	0.015	1.74	
Msn	Moesin	0.015	1.74	
Afdn	Afadin	0.021	1.70	
Col4a2	Collagen alpha-2(IV) chain; Canstatin	0.006	1.69	
Nomo1	Nodal modulator 1	0.014	1.68	
Col1a2	Collagen alpha-2(I) chain	0.0009	1.65	
Arpc4	Actin-related protein 2/3 complex subunit 4	0.007	1.62	
Sec23a	Protein transport protein Sec23A	0.030	1.59	
Patj	Pals 1-associated tight junction protein / InaD-like protein	0.003	1.53	
Lad1	Ladinin-1	0.012	1.49	
Rtn4	Reticulon-4	0.012	1.49	
Cisd2	CDGSH iron-sulfur domain-containing protein 2	0.026	1.46	
Naca	Nascent polypeptide-associated complex subunit alpha	0.049	1.46	
Scp2	Non-specific lipid-transfer protein	0.026	1.36	
Aimp1	Endothelial monocyte-activating polypeptide 2	0.046	1.26	
Dhrs7b	Dehydrogenase/reductase SDR family member 7B	0.024	1.24	
Clic1	Chloride intracellular channel protein 1	0.027	1.24	

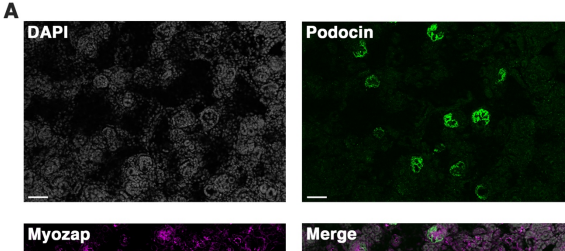
274 Supplemental Table V. Compiled proteomic profile of three cohorts of, 8-10 week old male and female, Nphs2<sup>BiolD2</sup> mice. Table list 54 significant proteins (p < 0.05) at a Log<sub>2</sub> cut off > 1.20, 275 with respective *p*-value listed in the far-right column. Table compiles the proteomic profiles across 276 all three separate analysis and averages their respective Log<sub>2</sub> fold change and *p*-values. This table 277 278 was subsequently utilized for Qiagen Ingenuity Pathway Analysis (IPA) and input into the Database for Annotation, Visualization, and Integrated Discovery (DAVID) for gene ontogeny 279 280 characterization. All proteins were detected with at least 2 razor unique peptides specific to the 281 protein denoted.

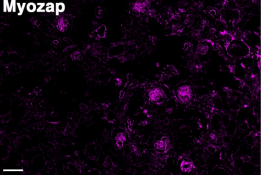
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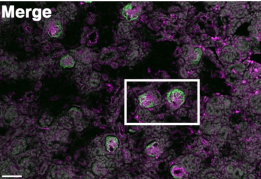
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Supplemental Figure 1. Expression of immunoglobulin superfamily member Jam1/F11r in tubule cells neighboring podocytes. (A). Jam1 was identified to have a 0.7 Log<sub>2</sub> fold change in a single mass spec analysis. P0 kidney sections were immunostained for Jam1 (magenta) and podocin (green). Jam1 is localized to tubule cells directly adjacent to podocytes. (A') Highlighted white boxed region from (A) is enlarged to show podocin localization restricted to the glomerulus, outlined in a dotted while circle, and Jam1 localization in neighboring tubule cells, denoted by white arrow. All scale bars: 50 μm.

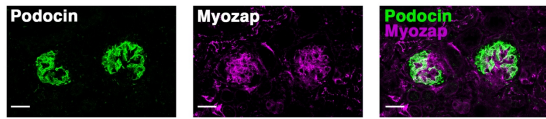
# Supplemental Figure 2.





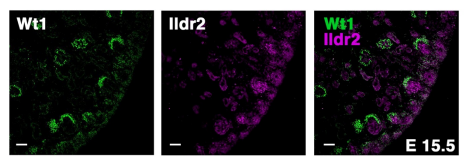






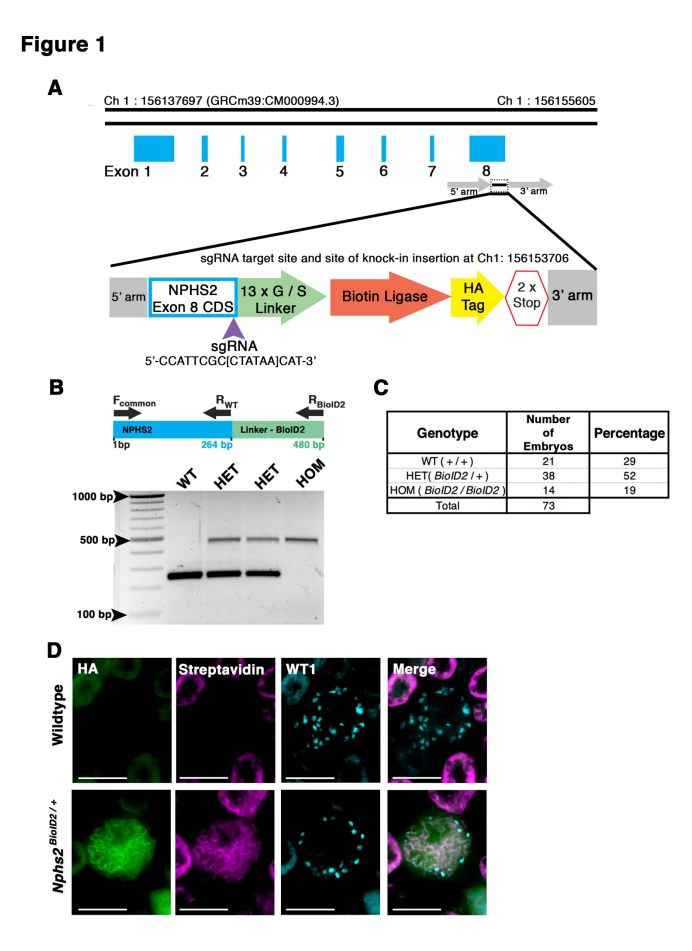
Supplemental 2. Myozap endothelium Figure localizes to of glomeruli. **(A)** Immunofluorescence analysis of Myozap (magenta) identifies strong localization to the central region of the glomerulus and endothelium outside the glomerulus. Proteomic profiling detected a significant Myozap signal (3.45 Log<sub>2</sub> FC), in two of three MS analyses. We observe some overlap (white) between Myozap (magenta) and podocin (green), while the majority of Myozap signal does not overalp with podocin and instead localizes to presumptive endothelium cells. Scale bar: 50 µm (A') Highlighted white boxed region from (A) indicating staining of Myozap within the endothelium of the glomerulus and some overlap with podocin. Scale bar 20  $\mu$ m.

# Supplemental Figure 3. A



**Supplemental Figure 3. Ildr2 is detected in early embryonic nephron precursors including the renal vesicle, comma, and s-shaped bodies.** IF analysis of E15.5 animals detects Ildr2 in both continuous membranous staining pattern and in punctate spots. In some S-shaped bodies, denoted by Wt1 expression in developing podocytes (green), Ildr2 (magenta) is detected in a continuous membranous pattern that appears to extend into the tubules, arrow. However, Ildr2 also appears punctate in other renal vesicles, comma, and s-shaped bodies, arrowheads.

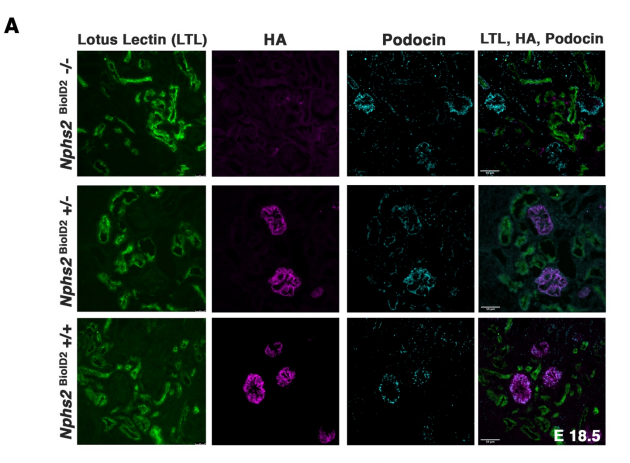
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- 2 Figure 1. Generation of a knock-in *Nphs2*<sup>BiolD2</sup> mouse line via CRISPR/Cas9 genome editing
- of the *Nphs2* locus. (A) Schematic of the CRISPR/Cas9 genome editing strategy utilized to
- 4 generate the *Nphs2<sup>BiolD2</sup>* mouse line. A small guide RNA (sgRNA) targeting the stop codon in exon
- 5 8 of *Nphs2* (purple arrowhead) was combined with a donor vector containing the knock-in cassette
- (zoom view) to induce homologous recombination and integrate the BioID2 moiety containing a
   13x Glycine/Serine (G/S) linker, biotin ligase, and HA tag into the *Nphs2* locus. (B) Genotyping
- strategy (top panel) to identify genotype of mice as wildtype (single band at 264 base pairs (bp)),
- 9 heterozygous (two bands; one at wildtype size of 264 bp and a second that amplifies the BioID2
- 10 linker region giving a band at 480 bp), or homozygous (single band at the 480 bp). (C) Genotyping
- 11 of 73 embryos at E18.5-P0, verify an approximate Mendelian ratio of genotypes being recovered
- 12 (25:50:25). (D) IF analysis of 8–10-week-old adult mice injected with biotin illustrate an enrichment
- 13 of streptavidin detected within the glomerulus of Nphs2<sup>BiolD2</sup> mice and absence of streptavidin
- 14 signal in control age matched C57BL/6J littermate mice. The HA signal from the BioID2 moiety
- 15 closely overlaps with streptavidin, observed as white overlap in the merged image. Wilms Tumor
- 16  $\,$  (WT1) is utilized as a podocyte marker to delineate the glomerular boundaries. Scale bar: 50  $\mu m$
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#### Figure 2.



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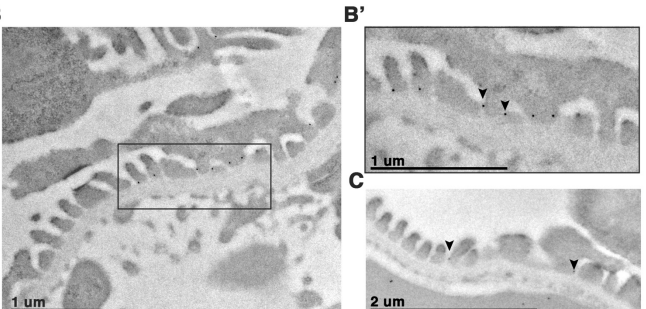


Figure 2. Nphs2<sup>BiolD2</sup> animals present normal kidney morphology with podocin-BiolD localized to podocytes and specifically the slit diaphragm. (A) E18.5 littermates of wildtype, heterozygous (Nphs2<sup>BiolD2/+</sup>), and homozygous (Nphs2<sup>BiolD2/BiolD2</sup>) animals were analyzed for gross kidney morphology and localization of podocin-BioID. The proximal tubule marker lotus lectin (LTL, green) show normal tubule architecture without observation of nephron tubule dilation in heterozygous (*Nphs2<sup>BiolD /+</sup>*), and homozygous (*Nphs2<sup>BiolD2 / BiolD2</sup>*) animals. The HA tag (magenta) which is part of the BioID2 moiety was observed to tightly overlap with podocin (cyan) specifically in the glomerulus. The relative size and number of glomeruli was unchanged between control, heterozygous, and homozygous animals. Further, the HA signal was not identified outside of the glomeruli. Scale bar 50 µm (B) Nphs2<sup>BiolD2/+</sup> localizes to the podocyte slit diaphragm. Utilizing transmission electron microscopy (TEM) and immunogold labeling of 4-week-old murine kidney samples, we detect the anti-HA signal as punctate dots strung just above the GBM adjacent to the podocyte foot process within electron dense regions where the slit diaphragm bridges neighboring foot processes. (B') Magnified view of boxed region from panel (B) depicting podocyte foot processes and the electron dense regions between them where the HA signal localizes as dark spherical dots from immunogold labeling, denoted with arrowheads. Scale bar 1 μm (C) Wildtype littermates were immunogold labeled for podocin and similarly display localization of podocin at electron dense regions between podocyte foot processes, highlighted with arrowheads. Scale bar 2 um.

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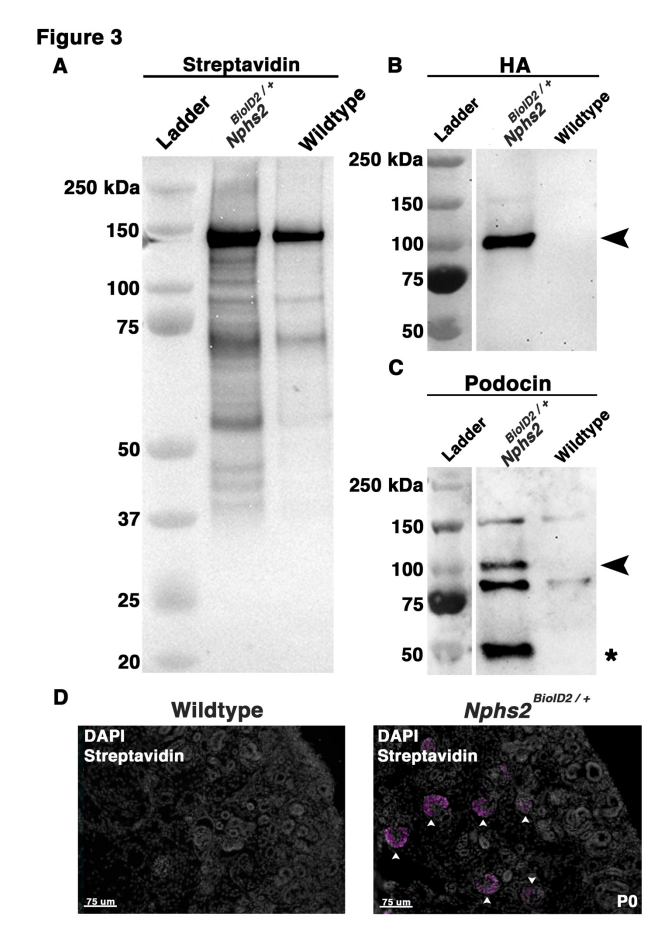


Figure 3. Biotin administered Nphs2<sup>BiolD2/+</sup> mice present an enrichment of biotinylated 72 proteins specifically within glomeruli. (A) A significant enrichment of streptavidin labeled, 73 74 biotinvlated, proteins are observed across the entire molecular weight spectrum in Nphs2<sup>BioID2/+</sup> 75 protein lysates compared to wildtype control littermates, injected with biotin. Streptavidin bead 76 purified lysates were subjected to protein separation and probed for streptavidin. Wildtype age matched, 8–10-week-old, littermate controls present few streptavidin conjugated bands, 77 presumably endogenous metabolic carboxylases. However, Nphs2<sup>BiolD2/+</sup> lysates exhibit numerous 78 specific biotinvlated protein bands across the entire molecular weight spectrum. (B) Nphs2<sup>BiolD/+</sup> 79 80 lysates blotted with an anti-HA antibody present robust and specific detection of the HA signal only within the Nphs2<sup>BiolD2/+</sup> sample, indicative of lysate purity. The HA signal is evident at approximately 81 100 kDa (arrowhead) in Nphs2<sup>BiolD2/+</sup> lysates. (C) Nphs2<sup>BiolD2/+</sup> samples probed for podocin show 82 83 two specific bands for both BioID2-tagged podocin and endogenous podocin. Adult 8-10-week-old Nphs2<sup>BiolD2/+</sup> mice blotted for podocin present a 50kDa band for the endogenous unmodified 84 85 podocin protein, marked with an asterisk (\*) and a second larger band at approximately 100 kDa for the HA tagged podocin-BioID of *Nphs2*<sup>BioID2/+</sup> mice (arrowhead). (D) P0 pups collected from 86 87 pregnant dams injected with 5 mg/kg of biotin every day from E11.5 to E18.5, and without injection of newborn pups, manifest strong streptavidin detection within glomeruli. Immunofluorescence 88 89 analysis reveals an enrichment of streptavidin signal (magenta) specifically within the glomeruli (arrowheads) of *Nphs2<sup>BiolD2/+</sup>* kidneys compared to wildtype. Scale bar 75 µm. 90

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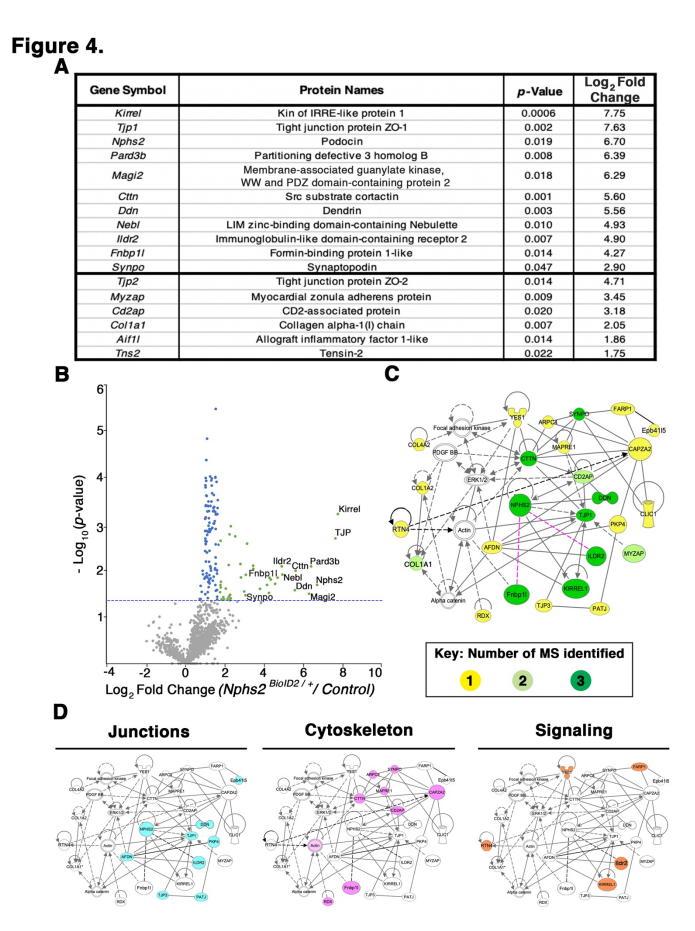
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106 Figure 4. Proteomics profiling of the podocyte foot process identifies documented slit diaphragm components and novel candidates with unexplored podocyte function. (A) Three 107 108 separate MS analyses were combined, and the average Log<sub>2</sub> fold change and respective p-values 109 were averaged to produce a list of the top foot process proteins. The top 11 proteins were 110 identified across all three MS profiles. Six additional proteins were identified across two of the three MS analyses, denoted following the thick black bar. Podocin was a top protein identified and 111 112 known to oligomerize, indicating our MS proteomic profiling was effective. Additionally, many of the 113 top proteins uncovered, *i.e.*, Kirrel, Dnd, Tjp1/2, Magi2, Pard3, Synpo, and Cd2ap are documented 114 podocyte slit diaphragm components. (B) A volcano plot depicting approximately 1400 proteins 115 identified across all three MS analyses. Green dots represent the 40 proteins identified across 116 three separate MS profiles as having a Log<sub>2</sub> fold change  $\geq$  1.7, and *p*-value  $\leq$  0.05, blue dots 117 denote all proteins identified to have a significant *p*-value < 0.05, and grey dots are proteins with a *p*-value > 0.05. The top 11 proteins consistently uncovered across all MS analyses are embedded 118 119 with gene symbols in the plot. The blue dotted line represents a p-value < 0.05. (C) Network topology was generated utilizing Qiagen Ingenuity Pathway Analysis (IPA) on the top 54 proteins 120 121 from our cumulative proteomic profiles. IPA produces a proposed web of relationships from 122 published literature with most of our top proteins represented, (11 of 17), with documented 123 connections to other foot process and slit diaphragm components. The color intensities of each protein (vellow to green) represent the number of MS analyses from which each protein was 124 identified. Darkest green shade=all three MS analyses, lighter green shade=two of three MS 125 analyses, and yellow=a single MS analysis. Dotted lines and unshaded proteins represent 126 127 predicted interactions / interactors from IPA analysis, respectively. Magenta lines are novel 128 proteins identified in this study to be present within the podocyte foot process. (D) Dissecting the 129 IPA network, identifies three central nodes representing junctions, cytoskeleton, and signaling. 130 Each respective node is color coded with junctions in cyan, cytoskeleton in magenta, and signaling in orange. These computational network topologies and nodes tightly align with podocyte structure 131 132 and function.

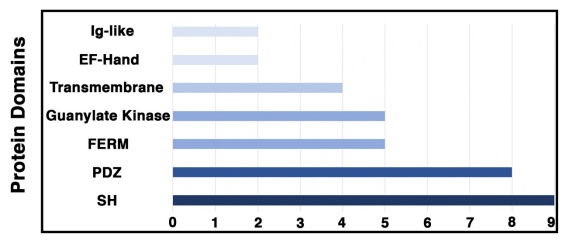
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### Figure 5.

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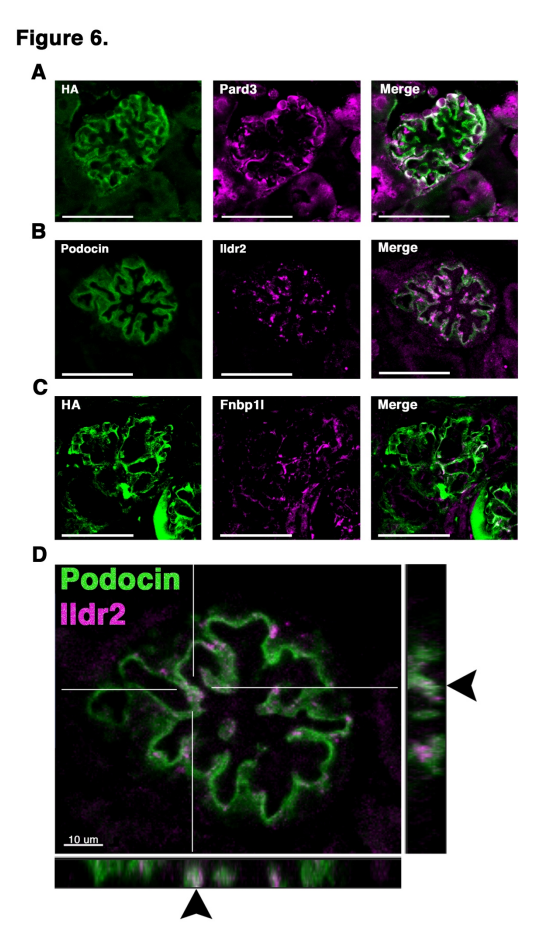
Gene Ontology (GO) Category	Cluster	Enrichment Score	Counts	p-Value
Molecular Function (MF)	Cytoskeletal Protein Binding	4.53	12	2.20 x 10
	Actin Binding	4.55	6	7.80 x 10
Cellular Component (CC)	Cell-Cell Junction	4.42	12	7.70 x 10
Cenular Component (CC)	Adherens Junction	4.42	10	7.90 x 10
Biological Process (BP)	Cytoskeleton Organization	2.61	14	3.20 x 10
Diological Flocess (BP)	Actin Filament-Based Process	2.01	12	8.70 x 10

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Number of Proteins w/ Domain from Foot Process Proteome

Figure 5. Analysis of the podocyte foot process proteome identifies the cytoskeleton, cell-135 cell junctions, and actin-based processes as the top GO categories with respective protein 136 137 domains that align with these functions. The Database for Annotation. Visualization, and 138 Integrative Discovery (DAVID) was utilized for GO clustering analysis of the top 54 proteins 139 identified to have a Log<sub>2</sub> fold change > 1.20, and p-value < 0.05 (Supplemental Table V). (A) Overlap in GO readout was observed across the three separate GO categories analyzed. 140 141 molecular function, cellular component, and biological process, with cytoskeleton, actin, and 142 cellular junctions being the top hits. The respective top clusters within each GO category are listed. 143 alongside the relative enrichment score, number of proteins, and *p*-values. (B) Protein domain 144 analysis of the top 54 podocyte foot process proteins provides evidence for protein-protein 145 interactions and likely scaffold and protein-protein complex formation. Each protein's respective 146 protein domains were binary counted for presence or absence within the proteomics profile. The top seven protein domains identified from the podocyte foot process proteome consist of a Src 147 148 homology (SH) (n = 9 proteins), PSD-95, Disc large, and ZO – 1 (PDZ) (n = 8), 4.1 protein, Ezrin, 149 Radixin and Moesin (FERM) (n = 5), Guanylate Kinase (n = 5), Transmembrane (n = 4), E and F 150 helix - Hand (EF - Hand) (n = 2), and an Immunoglobulin - like (lg - like) domain (n = 2). 151 Cumulatively, these 7 domains represent ~ 50 % of the top proteins identified from the foot process 152 proteome (26 / 54).



154 Figure 6. Two novel foot process candidates, Ildr2 and Fnbp1I, localize to podocytes and overlap with podocin in the foot process. IF analysis of three top candidates identified from the 155 proteomics profiling, Pard3, Ildr2, and Fnbp1I, were detected within glomeruli. (A) Pard3 156 (magenta), a known component of the podocyte foot process, co-localizes with HA signal of 157 158 podocin-BioID (green) producing a white signal in overlap. (B) Ildr2 (magenta) displays a punctate localization pattern within the glomerulus and overlaps (white) with podocin (green). (C) Fnbp11 159 160 (magenta) co-localizes (white) with HA signal of podocin-BioID (green) in podocytes. Scale bars in A-C: 50 µm (D) A merged image of podocin and Ildr2 through a confocal z-projection identifies 161 punctate localization and overlap (white, arrowheads) between podocin (green) and Ildr2 162 (magenta) visualized in the z-plane in the left and bottom panels. Scale bar in D 10  $\mu$ m. 163

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# Figure 7. A' A 南小田 Nphs2 B' В lldr2 C' С Fnbp1l P

166 Figure 7. In situ hybridization confirms glomerular and some tubule cell expression of Ildr2 and *Fnbp11* in P2 kidneys. P2 kidney sections were hybridized with antisense riboprobes against 167 Nphs2 (podocin), Ildr2, and Fnbp11. (A) Nphs2 (purple) displays strong expression specifically in 168 glomeruli at P2. The rectangular box in (A) is enlarged in (A') to highlight expression of Nphs2 only 169 170 in glomeruli, arrowheads denote example glomeruli (B) Ildr2 transcripts (purple) are identified within glomeruli and tubules. The rectangular box in (B) is enlarged in (B') to denote expression of 171 172 *Ildr2* in glomeruli, arrowhead, and in some tubules, black arrow. (C) *Fnbp11* (purple) is identified in 173 both glomeruli and tubules. The block rectangular box in (C) is enlarged in (C') to denote 174 expression of *Fnbp11* in glomeruli, arrowhead, and tubules, arrow. Scale bars in A-C: 500 μm. Scale bars in A'-C': 100 µm. 175

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# Figure 8.

Gene Symbol	Protein Names	BiolD (Young) vs WT Log <sub>2</sub> Fold Change	BioID (Aged) vs WT Log <sub>2</sub> Fold Change	<i>p</i> -Value
lldr2	Immunoglobulin-like domain-containing receptor 2	2.76	3.53	0.001
Tjp2	Tight junction protein ZO-2	2.71	3.31	0.01
Pkp4	Plakophilin-4	2.27	2.94	0.01
Nphs2	Podocin	2.42	2.36	0.10
Fnbp1l	Formin-binding protein 1-like	3.22	2.12	0.10
Tns2	Tensin-2	1.30	0.36	0.10

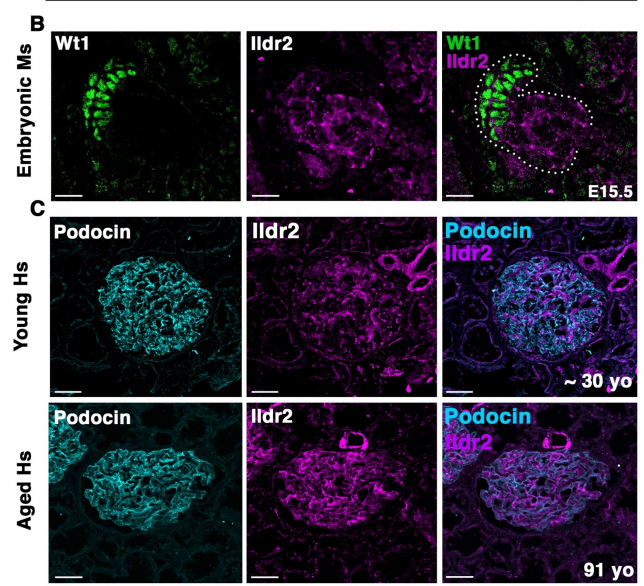
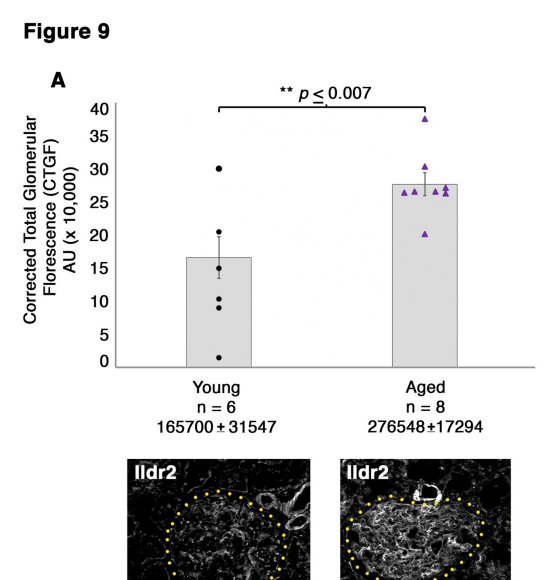


Figure 8. Ildr2 is detected in developing nephron structures and presents increased 183 detection in aged mouse and human glomeruli compared to young. (A) Proteomic profiling of 184 two aged (108 week old) male *Nphs2<sup>BiolD2/+</sup>* mice identifies a significant increase in Ildr2. Tip2, and 185 Pkp4 protein levels compared to 8-10 week old Nphs2<sup>BiolD2/+</sup> mice. Table lists 6 proteins with 186 187 relative detection in "young" 8-10 week old Nphs2<sup>BiolD2/+</sup> male mice vs "aged" 108 week old *Nphs2<sup>BiolD2/+</sup>* male mice. All protiens were detected with at least 2 razor unique peptides with their 188 189 respective *p*-values in far right column. (B) Ildr2 is expressed in early developing nephron 190 structures including the renal vesicle, comma, and s-shaped bodies with podocytes denoted by the 191 Wt1-positive nuclei. Ildr2 (magenta) is observed as membranous along the comma / s-shapped 192 body with some punctate foci in E15.5 mouse kidney cortex, outlined with white dots. Scale bar: 20 193 μm. (C) Young and aged (91 yo) human kidney tissue was probed of Ildr2 via IF. Young human 194 glomeruli display punctate foci detection of Ildr2 (magenta) similar to 8-10 week old mice. 195 However, aged (91 yo) male glomeruli evince an increase in Ildr2 detection within glomeruli. Furthermore, where Ildr2 looks to be more highly detected in these aged glomeruli podocin 196 197 appears less detectable. Scale bar: 50 µm.

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210 Figure 9. Aged 91 yo glomeruli exhibit a significant increase in corrected total glomerular florecence (CTGF) compared to young human glomeruli. We observe a significant (p < 0.007) 211 increase, of approximatly 67%, in CTFG in aged human Ildr2 detection compared to young human 212 tissue. CTGF was calculated by selecting the gomeruli as the ROI then measuring the total 213 214 florescence within the area and subtracting out the sum of the area by the average background florecence, *i.e.* Total gROI florecence – (Area of glomerulus x mean of background florescence) = 215 216 CTGF. Two example glomeruli are displayed and outlined in yellow dots to depict the respective ROI utilized to calculate CTGF. Scale bar: 50 µm. 217