On the Origin of Proteins in Human Drusen: The Meet, Greet and Stick Hypothesis

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1 **Title:** On the Origin of Proteins in Human Drusen: The Meet, Greet and Stick Hypothesis 2 3 **Abstract:** Retinal drusen formation is not only a clinical hallmark for the development of 4 age-related macular degeneration (AMD) but also for other disorders, such as 5 Alzheimer's disease and renal diseases. The initiation and growth of drusen is poorly 6 understood. Attention has focused on lipids and minerals, but relatively little is known 7 about the origin of drusen-associated proteins and how they are retained in the space 8 between the basal lamina of the retinal pigment epithelium and the inner collagenous 9 layer space (sub-RPE-BL space). While some authors suggested that drusen proteins are 10 mainly derived from cellular debris from processed photoreceptor outer segments and 11 the RPE, others suggest a choroidal cell or blood origin. 12 Here, we reviewed and supplement the existing literature on the molecular composition 13 of the retina/choroid complex, to gain a more complete understanding of the sources of 14 proteins in drusen. These "drusenomics" studies showed that a considerable proportion 15 of currently identified drusen proteins is uniquely originating from the blood. A smaller, 16 but still large fraction of drusen proteins comes from both blood and/or RPE. Only a 17 small proportion of drusen proteins is uniquely derived from the photoreceptors or 18 choroid. We next evaluated how drusen components may "meet, greet and stick" to each 19 other and/or to structures like hydroxyapatite spherules to form macroscopic deposits 20 in the sub-RPE-BL space. Finally, we discuss implications of our findings with respect to 21 the previously proposed homology between drusenogenesis in AMD and plaque 22 formation in atherosclerosis.

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1. Drusen.

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79 Drusen are extracellular deposits of bio-materials underneath the retinal pigment 80 epithelium (RPE) in the eye (Farkas et al., 1971b; Sarks, 1976). They are considered 81 clinical hallmarks for a number of diseases, including age-related macular degeneration 82 (AMD) (Hogan, 1965; Sarks, 1976; Hageman et al., 2001; Khan et al., 2016), Alzheimer 83 disease (AD) (Csincsik et al., 2018) and dense deposit disease (DDD) (Duvall-Young et 84 al., 1989; Mullins et al., 2000; Boon et al., 2009). AMD is the leading cause of severe 85 visual impairment, affecting 4% of the population over 60 years old (de Jong, 2006). AD 86 is the biggest cause of dementia, affecting millions of people in the western world. DDD 87 is a relatively rare juvenile disease characterized by kidney malfunction (Ito et al., 2017; 88 Wang et al., 2017; Cunningham and Kotagiri, 2018). Despite the potential relevance for 89 diseases, little is known about the composition of drusen and how and why biomaterials 90 accumulate these deposits. 91 Drusen are heterogeneous in terms of size, shape, color on retinal imaging, retinal 92 location and molecular content (Sarks, 1976; Sarks et al., 1980; Sarks et al., 1999; Crabb 93 et al., 2002; Khan et al., 2016). In the clinic, drusen can be identified as yellow spots on 94 funduscopy and color fundus images or dome shaped objects of different sizes under the 95 RPE on Optical Coherence Tomography (OCT) (Marshall et al., 1992; Bird et al., 1995; 96 Loeffler and Lee, 1998; Khan et al., 2016). Histopathological examination of drusen 97 showed that are located between the basal lamina of the RPE cells and the Inner 98 collagenous layer of the Bruch's membrane, a space that had been termed recently as 99 sub-RPE-BL space (Balaratnasingam et al., 2016; Li et al., 2018). Clinical definition of 100 drusen depends on size, color, auto fluorescence, and retinal location (Sarks, 1976; Bird 101 et al., 1995) (Figure 1). Drusen may appear in the macula, peri-macular area or in the 102 mid-and/or far periphery (Lengyel et al., 2015; Domalpally et al., 2017; Csincsik et al., 103 2018). A particular druse can be termed as "hard", when it's appearance is small, round and well demarcated, with a size of <63 μm . "Intermediate" drusen have a size of 104 105 approximately 63-125 μm, while "soft" drusen are >125 μm in size, and frequently have 106 more ill-defined edges (Bird et al., 1995). A few (<5) small hard (sub-clinical) drusen in 107 the macula does not raise alarm bells, but when numbers of hard drusen increase, or the 108 size of drusen increases such that they become "intermediate" and/or- "soft" drusen, the 109 likelihood to progression to AMD is increased significantly (Bird et al., 1995). Drusen

110 should be distinguished from reticular pseudodrusen (or subretinal drusenoid deposits) 111 that occur between the RPE and photoreceptor (PR) in the subretinal space (Zweifel et 112 al., 2010; Spaide et al., 2018). Relatively little is known about pseudo-drusen and as such, they are excluded from this review. Drusen are formed in the sub-RPE-BL space, 113 114 between the basement membrane of the RPE and the inner collagenous layer of Bruch's 115 membrane (BrM). 116 The RPE is a multifunctional single neuro-epithelial cell-layer that act as a metabolic 117 interface between the choroid and the neurosensory retina (Strauss, 2005). The RPE 118 cells are connected by intercellular tight junctions, together forming the outer blood-119 retina barrier. On the apical side, the photoreceptor cells line the RPE. On the basal side 120 the interposing BrM separates the basement membrane of the RPE from the choroidal 121 micro-vasculature (choriocapillaris). The choroidal capillaries are fenestrated, and not 122 surrounded by pericytes or smooth muscle cells. The BrM consists of three interleaved 123 layers: the inner and outer collagenous layers with an elastic layer in between them 124 (Booij et al., 2010a). Often, the basement membranes of the endothelium and the 125 epithelium are classified as part of the BrM but we will refer here to the BrM structure 126 as tri-laminar (rather than as penta-laminar). Embedded in the BrM are macromolecules 127 such as proteins and proteoglycans to help remodeling the extra cellular matrix (with 128 age)(Guo et al., 1999; Guymer et al., 1999; Del Priore et al., 2006; Beattie et al., 2010; 129 Booij et al., 2010a; Hussain et al., 2011). The diffuse thickening of BrM is also a 130 characteristic age-related feature (Hogan, 1965; Sarks et al., 1999). This is largely due to 131 the entrapment of proteins and lipids within the ECM (Curcio et al., 2011; Curcio and 132 Johnson, 2012). The diffuse build-up of extracellular bio-materials between the 133 basement membrane of the RPE and the inner collagenous layer of the BrM is called 134 basal linear deposits while the deposit formation between the basement membrane and 135 the cell membrane of the RPE are called basal laminar deposits (Sarks, 1976; Sarks et al., 1980; van der Schaft et al., 1993; Abdelsalam et al., 1999; Curcio and Millican, 1999; 136 137 Spraul et al., 1999). Due to the lack of information of the composition of these deposits, these specific classifications are excluded from our analysis. The deposits in BrM result 138 139 in a decline in the conductivity of the membrane creating in a diffusion barrier that 140 further enhances the accumulation of biomaterials (Green and Enger, 1993; Moore et al., 141 1995; Starita et al., 1997; Curcio and Millican, 1999; Curcio et al., 2011; Curcio, 2018b).

142 This phenomenon may be a general "passive" pathophysiological process that resembles 143 plaque formation in disorders such as AD or atherosclerosis. 144 Even more detailed insights into sub-RPE-BL space deposits originated from molecular 145 and histochemical studies on isolated drusen material. Recent investigations have 146 shown that drusen contain lipids, trace elements, including zinc, iron and calcium, as 147 well as a wide array of different proteins (Crabb et al., 2002; Curcio et al., 2011; 148 Thompson et al., 2015; van Leeuwen et al., 2018). The distribution of these components 149 is not uniform, neither within nor between drusen, further emphasizing the 150 heterogeneous nature of the deposits (Thompson et al., 2015). 151 Oxidative modification of lipids and proteins may result in the cross-linking of these 152 molecules and may contribute to deposit formation and drusenogenesis. Subsequently, 153 local cellular damage at the very early onset of AMD, via the complement cascade attack 154 on drusen compounds and the NLRP3 inflammasome (Edwards and Malek, 2007; Yuan 155 et al., 2010; Doyle et al., 2012), can lead to retinal damage and more advanced AMD. 156 Relatively few studies addressed the origin of proteins in the initiation and progression 157 of drusen (Mullins et al., 2000; Nordgaard et al., 2006; Cryan and O'Brien, 2008; Wang et 158 al., 2010; Crabb, 2014). A number of studies (Johnson et al., 2011; Kunchithapautham et 159 al., 2014) have yielded conflicting data as to where drusen proteins originate from, and 160 whether the accumulation of this apparent depositioning of biomaterials in BrM is a 161 passive or an active process. Several questions remain, which include: to what extent do 162 proteins in drusen originate from photoreceptors, RPE, choroidal endothelium or even 163 the circulating blood? How do drusen form and how are drusen components recruited 164 and deposited in the sub-RPE-BL space? What is the extent of the (molecular) 165 heterogeneity that exists within and between drusen? Here, we will review and combine data from the existing literature, and supplement these with our own (new and recently 166 167 published) data from subretinal transcriptomic, proteomic and immunohistochemical 168 staining experiments. To enable this, we have functionally annotated a compiled list of 169 drusen proteins and compared these proteins with those identified in specific 170 transcriptomic and proteomic datasets derived from cells and tissues of the various 171 relevant compartments. These include both subretinal and choroidal tissues, as well as 172 the plasma proteome. Collectively, these analyses increase our understanding of 173 drusenogenesis, which may provide clues for the prevention of drusen formation and, 174 ultimately, for the prevention of drusen associated disorders (Khan et al., 2016)

2. Functional annotation of drusen proteins.

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177 One of the key aims of this study is to identify the most likely original sources of drusen 178 proteins. More specifically, do drusen proteins only come from the neural tissues 179 (photoreceptors and RPE) or is there also a choroidal or systemic component? In the 180 next chapters, we try to answer this question through a literature search and by using a 181 variety of qualitative and quantitative transcriptomics and proteomics meta-analyses of 182 the relevant genes and proteins involved. 183 We did not distinguish between various drusen types, sizes and/or drusen locations, 184 since little -omics data are available for each drusen subtype. Essentially, we followed 185 the (sub-clinical) drusen type description used by Crabb and coworkers (Crabb et al., 186 2002) who defined drusen to appear as opaque, 0 to 250 µm spherical to irregular 187 deposits that remained attached to BrM after removing the RPE from human donor 188 globes, both in the macular and the retinal periphery. 189 Based on relevant studies in the literature (Mullins et al., 2000; Crabb et al., 2002; Wang 190 et al., 2010), we curated a list of 89 drusen proteins (Table 1). This was achieved by 191 combining the published datasets and removing incomplete, duplicate or ambiguous 192 entries. Several entries did not correspond to a single full-length cDNA annotated in the 193 knowledge database Ingenuity (www.ingenuity.com) and were left out. Since the 194 complement gene pathway is likely the best and most extensively studied pathway 195 (compared to other pathways) we only added a few complement proteins to the list, to 196 avoid bias toward one pathway and the "winner's curse". In addition, we also searched 197 the literature for confirmatory immunohistochemistry (IHC) studies and manually 198 added proteins from such smaller-scale studies. We realize this list may not be complete. 199 For example, individual entries like the locally produced vitronectin (Hageman et al., 200 1999; Wasmuth et al., 2009) present in drusen is missing in Table 1 and an entry like 201 elastin may be present as contamination of the BrM rather than a "specific" drusen 202 protein. The problem with selecting these proteins lies with the heterogeneity of drusen 203 (one protein may be present in one drusen but not in the other), lack of uniform criteria 204 "what is drusen-specific (?)", lack of uniformity in healthy or diseased stage of examined 205 samples and overall, how much evidence is needed to assign proteins to drusen (see also

discussion section). Nonetheless, we believe that, for the purposes of this study, our

208 (Crabb et al., 2002) provides us with a sufficient representative drusen protein dataset 209 for the purpose of this study. 210 We used the 89 drusen protein data set first to investigate the molecular aggregation 211 and the functional annotation of drusen proteins. A similar study was previously carried out by Crabb and coworkers (Crabb et al., 2002; Crabb, 2014). However, here we used a 212 213 slightly different list of drusen proteins and subjected this to additional, advanced 214 bioinformatics analysis. Consequently, we ran an Ingenuity knowledge database core 215 analysis using our list of drusen components (Table 1) which yielded biological motifs, 216 canonical pathways and molecular structural or functional networks. A summary of the 217 results of this analysis is shown in Table 2. 218 2.1. Biological or disease motifs and canonical pathways. 219 The functional annotation of the 89 drusen proteins (Table 2), revealed that these proteins (motifs or aggregates) can be associated with a number of functional or disease 220 221 entities, such as "hereditary disorders", "ophthalmic disease", "organismal injury or 222 abnormalities" and "metabolic disease and developmental disorders". Although these 223 annotation categories are broad and not very specific, they do point to a wide range of 224 potential sources of drusen components from both local and systemic origin. 225 Ingenuity analysis also yielded a number of canonical pathways. A canonical pathway is 226 the simplest linear representation of an established chain of biochemically related 227 molecules in a given system or cellular environment. The software recognizes enriched 228 canonical pathways specific for "acute phase response signaling", the "retinoid- and 229 farnesoid X receptors (LXR/RXR and FXR/RXR) response", "atherosclerosis signaling" 230 and "IL-12 signaling in macrophages" in the drusen dataset. 231 "The acute phase response" is a fast-systemic inflammatory response triggered by 232 infection, tissue injury and/or immunological disease (Serhan et al., 2015). The response 233 is mediated by the hypothalamus and several acute phase plasma proteins. These 234 proteins have a broad-working spectrum: they kill micro-organisms and modulate 235 complement activation, enzyme activity and the immune response. How and why these 236 proteins potentially end up in drusen is not clear (Johnson et al., 2000). Although not 237 undisputed, Despriet and coworkers found independently, that AMD is associated with

selection of 89 proteins, largely based on the proteomic study of Crabb and colleagues

238 acute phase plasma protein levels and with genetic variation in C-reactive protein (CRP), 239 one of the principal acute phase proteins (Despriet et al., 2006). Chirco and Potempa 240 showed that CRP protein acts as a mediator of complement activation and inflammatory 241 signaling in AMD (Chirco and Potempa, 2018). It is generally assumed that acute phase 242 proteins are present in the blood; suggesting that some drusen proteins can originate 243 from this pathway and have a systemic origin. While choroidal CRP apparently correlate 244 to serum levels (Chirco et al., 2018), it cannot be said with certainty that these proteins 245 are not (transiently and/or locally) produced by the choroid as well in cases of (nearby) 246 low-grade inflammation. 247 "The retinoid X receptors (RXRs)" are nuclear retinoid receptors that regulate, via the 248 ligand LXR, lipid and cholesterol metabolism as well as inflammation (Hiebl et al., 2018). 249 Cholesterol metabolism is essential for many retinal functions (Pikuleva and Curcio, 250 2014), while ocular (para-) inflammation is crucial for maintaining retinal homeostasis 251 (Xu et al., 2009). In the eye, retinoid X receptor activation contributes to retinal 252 photoreceptor differentiation, survival, and disease (Forrest and Swaroop, 2012), and 253 more specifically, for docosahexaenoic acid-mediated protection of photoreceptors 254 (German et al., 2013). The presence of this protein signature in drusen points toward a 255 local cellular origin of this protein. LXR can form heterodimers with "the farnesoid X 256 receptor (FXR)" which is also a nuclear receptor, and is an important regulator of a 257 variety of bile acid, glucose and lipid-related metabolic pathways, including the removal 258 of cholesterol (Tu et al., 2000; Hiebl et al., 2018). FXR protein was detected in a variety 259 of tissues, including heart, ovary, thymus and eye. Both LXR and FXR may be involved in 260 cholesterol homeostasis in RPE and retina (Zheng et al., 2015). The presence of these 261 receptor proteins in drusen points toward a local cellular origin. 262 "Atherosclerosis signaling": Atherosclerosis is a low grade chronic inflammatory 263 disorder characterized by local plaque deposition in the vessel wall, formed by a local 264 accumulation of modified plasma lipoproteins and macrophage activation. The major 265 cause of coronary events is rupture and thrombosis. Interestingly, clinical, 266 epidemiological, pathobiological and molecular evidence suggest that an overlap exists 267 between drusen in AMD and plaque formation in atherosclerosis. Indeed, like AMD, 268 atherosclerosis is now considered as a low-grade chronic inflammatory process 269 resulting from interaction (in) between plasma lipoproteins and the vascular wall 270 (Mullins et al., 2000). In AMD, not only plasma lipoproteins, but also local lipoproteins

271 are involved. In section 8 of this manuscript, we describe the potential molecular and 272 pathobiological overlap between drusen/AMD and vascular plaques in detail. Taken 273 together, the homology between drusen and atherosclerotic plaques points toward a 274 systemic origin of some drusen proteins. 275 "IL-12 Signaling and Production in Macrophages": The production of the cytokine IL-12 276 by activated (incoming) macrophages in damaged or diseased retinal tissue, is well 277 known (Zamiri et al., 2006; Chen et al., 2013). However, IL-12 exerts an autocrine effect 278 since macrophages and dendritic cells also respond to IL-12 by producing interferons 279 that stimulates T-helper cell differentiation. The RPE is apparently able to suppress 280 inflammation by modulating IL-12 production (Zamiri et al., 2006). Cao and coworkers 281 showed that cultured RPE cell in vitro secrete several cytokines, including IL-12, under 282 conditions of oxidative stress and replicative senescence (Cao et al., 2013). Therefore, 283 the molecules identified in this category (IL-12 signaling) can originate from both the 284 circulation as well as from the local cellular environment. 285 2.2. Molecular networks. 286 Molecular networks in Ingenuity are built up from a myriad of relevant literature 287 connections and they are formed on the basis of most likely physical or functional 288 interactions between (input) genes and/or proteins. For example, see molecular 289 network 1 in Figure 2a. Based on millions of experimentally verified and curated data 290 points, these networks represent the most likely functional associations between 291 components of the "biological soup" in the context of the input molecules. Structural, 292 functional and mixed molecular networks exist. Structural networks contain primarily 293 networks of structurally and physically interacting entries. Functional networks are 294 dominated by functional relationship between participating molecules. A third, "mixed" 295 network, contains both structural and functional associations. The molecular network 296 analysis of drusen proteins yielded 4 significant networks, with 6 distinct functional 297 clusters. Note that the networks are not *a priori* built through their possible relationship

299 2.2.1. Network 1.1, 1.2 and 1.3: Complement, collagens and crystallins.

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with drusen or AMD *per se*.

The most significant network formed in the data-driven Ingenuity drusen analysis, is presented in Figure 2a. This network consists of three functionally more or less specific molecular clusters: the complement protein cluster, the collagen protein cluster and the

303 crystallin heat shock protein cluster (Table 3). The presence of complement proteins in 304 drusen (and the choriocapillaris) was previously shown in older and AMD affected eyes 305 through immunohistochemistry, long before the genetic involvement of CFH and other 306 complement factors in drusen formation and AMD became genetically apparent 307 (Johnson et al., 2000; Hageman et al., 2001; Edwards et al., 2005; Hageman et al., 2005; 308 Haines et al., 2005; Klein et al., 2005). Analyzing the drusen proteome, we confirmed the 309 involvement of the terminal complement protein complex by identifying the 310 complement factors C7, C8A, C8B, C8G, and the membrane attack complex (MAC) in 311 drusen. Of note, the MAC was initially identified in drusen from unspecified retinal 312 locations, but was later shown not to be present in macular drusen (Johnson et al., 2000; 313 Mullins et al., 2014). The MAC is the final downstream event of the complement cascade. 314 It results from the binding of C5b to blood plasma complement proteins C6, C7, C8, and 315 C9, forming transmembrane pores that leads to cell lysis and death. In the same cluster, 316 we found the Prolyl endopeptidase-like protein (PRELP), a small leucine-rich 317 proteoglycan (SLRP) (Hultgardh-Nilsson et al., 2015), which among others, is involved in 318 the inhibition of complement activation (Warwick et al., 2014). The main complement 319 cascade regulator CFH is another member of the complement cascade that is present in 320 the drusen proteome. Genetic variation in CFH may regulate complement activation on 321 RPE cells (Radu et al., 2014). Of note, is that a certain degree of low-grade complement 322 activation and para-inflammation is always present in healthy aging eyes, to maintain 323 local health (Xu et al., 2009). In a recent review, Warwick et al. concluded that 324 complement deposition in the retina could be of local and/or systemic origin (Warwick 325 et al., 2014). The majority of complement genes are expressed in the liver, resulting in 326 an abundance of complement proteins in the blood. However, the RPE expresses several 327 key complement genes which may modulate the complement attack on RPE and drusen 328 (Chen et al., 2007; Kim et al., 2009; Pao et al., 2018). Interestingly, locally produced CFH 329 is, at least in cultured RPE cells, secreted apically, and not basally (Kim et al., 2009; Pao 330 et al., 2018). Consequently, the potential regulating role of locally produced CFH in vivo, 331 and other complement factors, potentially involved in the complement attack on drusen, 332 needs to be further investigated. A diversity of collagen proteins, such as COLA1, A2, 333 6A1, 6A2 and 8A1 were previously consistently identified in basal laminar deposits and 334 basal linear deposits, and, occasionally, in drusen (Newsome et al., 1987; Booij et al., 335 2010a; Curcio and Johnson, 2012). Newsome and coworkers (1987) noted that the

336	involvement of extracellular matrix components in drusen is variable but these findings
337	have not been confirmed in other studies. These molecules may primarily present as a
338	remnant from the (ab)normal turnover of BrM components (Newsome et al., 1987).
339	Alternatively, they may be secreted by the RPE in response to challenges presented by
340	drusen or by the conditions that lead to drusen formation. Interestingly, collagen IV is
341	not present in our curated drusen dataset, despite the fact that collagen IV
342	accumulations are found in autosomal dominant radiant drusen in Doyne Honeycomb
343	Retinal Dystrophy caused by
344	EFEMP1(EGF-containing fibulin extracellular matrix protein 1) mutations (Sohn et al.,
345	2015). In fact, in the absence of detailed electron microscopic examination it is not clear
346	whether these are drusen or basal laminal deposits.
347	The presence of crystallin proteins in drusen had been shown by Crabb and co-workers
348	(Crabb et al., 2002) and functionally studied by Nakata et al. (Nakata et al., 2005). These
349	authors found that BrM, drusen and part of the choroidal connective tissue, when
350	affected by AMD, showed higher immunoreactivity for $\alpha\text{-}$ and $\beta\text{-}crystallins$ than healthy
351	control tissues. Retinal crystallins are also up-regulated in a variety of other retinal
352	pathologies, including diabetic retinopathy, ischemia, mechanical injury and uveitis. The
353	$\alpha\text{-crystallin}$ family plays a crucial role in neuroprotection and inflammation (Fort and
354	Lampi, 2011), while the $\beta\text{-}$ and $\gamma\text{-}crystallins$ are small proteins with a possible ganglion
355	cell protective role in glaucoma (Anders et al., 2017) and a role in retinal tissue
356	remodeling and repair (Thanos et al., 2014). Consequently, the presence of these
357	proteins in drusen points to a local cellular origin.
358	2.2.2. Network 2.4 development, genetics of ophthalmic disorders.
359	The fourth cluster in our drusen protein analysis is actually similar to the entire network
360	2 which is functionally annotated as "network of genetic and developmental disorders".
361	The components of this cluster are functionally presented in detail in Figure 2b; Table 3.
362	This network contains annexin A2 (ANXA2), a relatively small calcium and
363	phospholipid-binding protein involved in multiple intra-cellular transport functions. The
364	RPE secretion data set, called RPE-IVS (Table 3; STable 1) reveals that this protein is
365	indeed secreted basally by the RPE (Pao et al., 2018). The protein was initially assigned
366	to drusen (Crabb et al., 2002). However, the same authors showed, using IHC in a

- number of human donor eyes, that ANXA2 is not associated with the interior of drusen,
- but with the basal lamina of the RPE close to the drusen surface.
- 369 2.2.3. Network 3.5: Immunological response.
- 370 The fifth functional cluster in our drusen protein analysis represents network three:
- 371 "Injury and inflammatory response; dermatological disease" (Figure 2c). The
- components are given in Table 3. This network contains, among others, annexin A1
- 373 (ANXA1). ANXA1 antibodies intensely stained whole drusen, but also the BrM and
- 374 choroid (Rayborn et al., 2006). Given its positive staining in entire drusen, we consider it
- here as a drusen protein. Apolipoprotein E (APOE) is also present in this group. APOE is
- 376 classically thought of as a cholesterol carrier. Risk alleles of the APOE gene were
- 377 associated with a variety of diseases including AMD, AD and atherosclerosis (Klaver et
- 378 al., 1998; Ashford, 2004; Song et al., 2004; Tikellis et al., 2007). Its presence in laminar
- deposits and drusen was initially established by Klaver and colleagues (Klaver et al.,
- 380 1998) and later confirmed by Anderson and Malek (Anderson et al., 2001; Malek et al.,
- 381 2003). Interestingly, in an RPE cell culture model that mimics drusen formation, Pao
- 382 (2018) and coworkers found that APOE is secreted basally by these cells. Subsequent
- 383 exposure of these cultures to human serum led to heterogeneous sub-RPE-BL space
- deposits, some of which were rich in serum-derived proteins such as vimentin, clusterin
- and amyloid P (Pao et al., 2018). In addition to ANXA1 and APOE, the serum amyloid
- proteins S100A7, S100A8 and S100A9 are part of this functional cluster and the drusen
- proteome. S100 proteins are a family of small calcium-binding proteins, produced in the
- nucleus and cytoplasm of a wide variety of cells (Gross et al., 2014; Narumi et al., 2015;
- 389 Cunden et al., 2017).
- 390 2.2.4. Network 4.6 Cell-to-cell signaling and systemic involvement, lipid metabolism.
- 391 The sixth cluster "cell-to cell signaling and systemic involvements" (Figure 2d; Table 3)
- 392 points to proteins which come from an extracellular environment. For example, The
- 393 APOA1, APOA4 and SAA1 lipoproteins and S, and the protein-groups related to the LDL,
- 394 HDL, VLDL metabolism (that have been added by Ingenuity to construct a meaningful
- network) are most likely derived from the blood, and not from the retina. However,
- 396 cautious interpretation of these general data is warranted, since the RPE is also capable
- of secreting a number of lipoproteins, such as APOB (Li et al., 2005b). The mechanisms

398 of biogenesis of lipid-laden soft drusen has been recently reviewed elsewhere (Curcio, 399 2018a, b) as has the role of lipids in AMD (van Leeuwen et al., 2018). 400 In at least two blood proteomics datasets (Table 3) the ORM1 (acute phase plasma 401 protein of unknown function; www.genecards.org) and the SERPINA1 (serine protease 402 inhibitor; www.genecards.org) proteins occur, which point also at a systemic origin of 403 these drusen proteins. Furthermore, in this cluster we see the drusen protein clusterin 404 (CLU), which is expressed in many cell types, including photoreceptors or RPE, and is 405 also present in blood (Garcia-Aranda et al., 2018). The presence of annexin 6 (ANXA6) in 406 drusen (and BrM) was previously confirmed using immunohistochemistry (Rayborn et 407 al., 2006). Finally, we observe also the presence of the (systemic) HRG protein, which is 408 extensively discussed in section 7 of this manuscript.

3. Drusenomics, part I: Where do drusen proteins come from: *the literature*.

410 Multiple epidemiological, genetic, biochemical and pathophysiological studies in the 411 literature address the origin of drusen. While many studies address the origins of metal 412 ions or lipids in drusen, here we focus on the likely source of proteins. Drusen proteins 413 could originate from either the neural side of drusen (Photoreceptors, RPE), the 414 systemic side (BrM, choroid complex, blood) BrM, or both (Penfold et al., 2001; Curcio 415 and Johnson, 2012). 416 3.1. The neural side of drusen. 417 Theories on drusen accumulation from the neural side vary: proteins may either come 418 from dying PR and RPE cells, or from (basal) secretion of proteins generated by the 419 normal functions of the RPE (Crabb et al., 2002; Kinnunen et al., 2012). Respectively, 420 cellular debris or secreted proteins may get trapped in BrM or drusen. Evidence for 421 these origins was gathered from histopathological investigation, retinal imaging, and 422 proteomics studies. 423 3.1.1. Histopathological and retinal imaging observations. 424 Drusen formation goes hand in hand with hypo- or hyperpigmentation (Curcio et al., 425 1998) of the RPE, especially in the early stages of AMD. Indeed, retinal cells overlying 426 drusen exhibit numerous irregular structural and molecular abnormalities which are confined to areas directly internal to drusen (Farkas et al., 1971b; Hogan, 1972; Burns 427 428 and Feeney-Burns, 1980; The Eye Disease Case-Control Study, 1992; Johnson et al., 429 2003). Deflection and shortening of rod inner and outer segments of rod photoreceptors 430 have been postulated to contribute to sub-RPE deposit formation (Farkas et al., 1971a). 431 Drusen have been also associated with more indirect changes, such as alterations in the 432 synaptic terminals of photoreceptor cells and an increase in vimentin and glial fibrillary 433 acidic (GFAP) protein within Müller cells (Johnson et al., 2003). Other retinal cells, such 434 as bipolar, horizontal, amacrine and ganglion cells are most likely unaffected by 435 drusenogenesis (Johnson et al., 2003). 436 Using immunohistochemical, molecular biological and biochemical approaches, 437 Hageman and coworkers found that RPE cell loss is correlated with increasing drusen 438 density (Hageman et al., 2001). More recent OCT studies, focusing on the integrity of the

RPE layer directly internal to drusen showed that 41.3% of all drusen coincided with an

440 intact overlying RPE, and that in 28.1% of cases, the RPE was irregular but continuous 441 (Schlanitz et al., 2018). In 30.6% of cases, the RPE layer adjacent to drusen was 442 discontinuous. Larger drusen were associated with higher probability of RPE loss 443 (Schlanitz et al., 2018). Taken together, these results suggest that RPE or PR cell death is 444 associated with drusenogenesis. However, it is not clear whether the observed cellular 445 damage is a cause or consequence of sub-RPE deposit formation. 446 The presence of cytoplasmic (Burns and Feeney-Burns, 1980), fibrous and 447 membranous/lipoid material (Fine, 1981; Young, 1987; Green and Enger, 1993; Loeffler 448 and Lee, 1998; Curcio and Millican, 1999) in drusen suggest that deposits are formed 449 after cellular degeneration. According to Coats, small colloid bodies derived from 450 degenerated RPE cells, develop into larger drusen due to uptake of biomolecules 451 through a defective BrM (Coats, 1905) and clinical support was provided for the 452 existence of these bodies (Pauleikhoff et al., 1990). Later, necrotic RPE cells were 453 presumed to be incorporated into existing drusen (Young, 1987). However, these 454 findings also did not distinguish between cause or consequence of deposit formation. To 455 complicate matters further, there are a number of reports in the literature describing 456 drusen regression; in an experimental study after laser photocoagulation and in clinical 457 studies using fluorescein angiograms (FAs) fundus photography (Bressler et al., 1995) 458 and OCT (Yehoshua et al., 2011). A similar observation were done in rhesus monkeys 459 (Duvall and Tso, 1985) in APOE mice with thickened BrM as well as AMD patients 460 (Jobling et al., 2015). This intriguing phenomenon may be linked to transiently 461 increasing the RPE-mediated release of active MMP enzymes that alter the turnover of 462 BrM (Zhang et al., 2012). 463 3.1.2 Proteomic level observations. 464 Proteomics studies into drusenogenesis can be divided into studies on (archived) 465 human post-mortem eyes, in vitro RPE culture, and proteomic studies on retinas of 466 animal models. A variety of techniques, such as 2D gels and LC-MS/MS analysis have 467 been used. To date, up to over 500 healthy and AMD-affected post-mortem human eye 468 tissue specimens (numerous contributions of Sarks, Hageman, Mullins, Lutty, Bergen, 469 Lengyel, and Curcio) have been examined by light, confocal, or electron microscopy, in 470 conjunction with proteomics and with antibodies to specific drusen-associated proteins 471 (Curcio et al., 2017). These studies emphasize the heterogeneity of drusen, a concept

472	initially developed by Sarks and coworkers (Sarks et al., 1980; Sarks et al., 1994; Sarks et
473	al., 1999) and strongly suggest that chronic local inflammation at the level of BrM is an
474	important contributor to drusenogenesis.
475	<i>In vitro</i> , the transcriptome and proteome of RPE cells, such as cultured primary retinal
476	cells (fetal or from postmortem human donor eyes) (Alge et al., 2003; Oshikawa et al.,
477	2011; Pao et al., 2018) has been determined. Stable isotope labeling of amino acids
478	showed that these cells secrete a variety of extracellular matrix proteins, complement
479	factors, and protease inhibitors, that have also been reported to be major constituents of
480	drusen (An et al., 2006). In addition, abnormal protein secretion by human primary RPE
481	cultures derived from AMD patients has been observed compared to age-matched
482	controls (An et al., 2006). However, the fact that major components of drusen can be
483	reproduced by RPE cells without the need for PR outer segments, supports a crucial role
484	of RPE in drusen formation (Pilgrim et al., 2017). At the same time, it suggests that PRs
485	may contribute but are not essential for drusenogenesis. Off note, it is important to
486	emphasize that cells in culture were treated with heat-inactivated serum, and that the
487	contribution of components from this material to drusenogenesis, as "dietary"
488	contribution, is highly likely (Bretillon et al., 2008; Pikuleva and Curcio, 2014; Pilgrim et
489	al., 2017).
490	Wang and coworkers found that, after simultaneous mass spectrometry analysis of both
491	archived drusen and RPE material, similar protein profiles, but with higher intensities
492	and greater variability in the drusen. Within the limits of unavoidable sample
493	contamination, these data suggest that other than RPE alone, additional local cells or
494	tissues contribute to formation of debris in the sub-RPE-BL space (Wang et al., 2010).
495	3.2. The systemic side of drusen.
496	Drusenogenesis theories have focused on the role of lipids and immune-mediated
497	effects. Lipoproteins, neutral lipids (Curcio et al., 2011), complement-activating
498	molecules and other immune mediators as well as monocyte-derived cellular processes
499	have been identified within drusen (Hageman et al., 2001; Penfold et al., 2001; Anderson
500	et al., 2010; Molins et al., 2018), which indicates the biogenesis or propagation of drusen
501	from the systemic side.

3.2.1. Bruch's membrane.

503 The main functions of BrM are structural, to support the RPE, and to regulate the 504 transport of fluid, ions and biomolecules from the choroid to the RPE, and vice versa 505 (Curcio and Johnson, 2012). BrM thickening and decline of hydraulic conductivity have 506 been observed during aging (Hussain et al., 2010; Cankova et al., 2011). Studies suggest 507 diffuse thickening of the inner aspect of BrM is associated with retinal pigment epithelial 508 hypopigmentation, focal atrophy, and soft (large) drusen formation (Bressler et al., 509 1994). A variety of extracellular matrix components have been detected in diffuse 510 thickenings of BrM (Fernandez-Godino et al., 2016). Immunohistochemical reactivity of 511 BrM showed age-related accumulation of type I collagen and localized changes 512 associated with some drusen (Newsome et al., 1987; Curcio and Johnson, 2012). The 513 tissue inhibitor of metalloproteinases-3 (TIMP-3) protein, a major component of the 514 drusen proteome, showed high immune-reactivity in human drusen and in BrM (Fariss 515 et al., 1997). The continuous turnover of BrM during life could provide a continuous 516 local supply of BrM proteins. Some of the remnants may be cleared to the blood but 517 some of them might end up in drusen. Please note, that most studies on the aspects of 518 BrM thickening have been performed by light microscopy on paraffin sections. In future 519 studies, it will require TEM or high resolution light microscopy to confirm the majority 520 of these findings, and to distinguish, for example, between "BrM thickening" and basal 521 laminar deposits. 522 3.2.2. Choroidal capillaries. 523 The choriocapillaris is located directly underneath the RPE and BrM. It is composed of a 524 unique vascular network which provides nutrients and fluid for the RPE and the retina 525 (Bernstein and Hollenberg, 1965). The abundance of fenestrations on the RPE aspect of 526 the choriocapillaris endothelium makes this vascular bed much leakier than non-527 fenestrated vessels (Bernstein and Hollenberg, 1965). A compromised interface can 528 result in various abnormalities such as choroidal neovascularization (CNV) and AMD 529 (Lutty et al., 2010). 530 With age and in AMD, the choroid thins. The choriocapillaris loses density and covers an 531 increasingly smaller portion of BrM. At the same time, increased drusen deposition 532 occurs, as witnessed by histopathological evidence (Ramrattan et al., 1994; Ida et al., 533 2004). OCT Angiography (OCTA) showed atrophy of choriocapillaris underneath and

beyond the region of photoreceptors and RPE loss (Wakatsuki et al., 2015; Moreira-Neto

535 et al., 2018), in agreement with previous and parallel histopathological studies (McLeod 536 et al., 2009; Biesemeier et al., 2014). In human macular sections, histopathological 537 evaluation of the sub-RPE-BL deposits together with potential vascular changes, showed 538 that vascular density was inversely correlated with sub-RPE-BL deposit density 539 (Biesemeier et al., 2014). Curcio and coworkers observed that modest endothelial cell 540 loss in the choriocapillaris also occurred directly adjacent to basal linear deposits and 541 subretinal drusenoid deposits (Curcio et al., 2013). Sub-RPE-BL deposits showed a 542 positive correlation with the number of ghost vessels in the choroid, suggesting that 543 vascular endothelial cell loss could contribute to deposit formation (Mullins et al., 2011). 544 It has also been shown that the presence of complement components and specifically, 545 MAC, in the choroid increases with aging, and increases even more in AMD-affected eyes 546 (Mullins et al., 2014; Chirco et al., 2016). In fact, C5b-9 complement complexes are present in hard drusen, BrM, and extend to the choriocapillaris in some cases (Johnson 547 548 et al., 2000; Anderson et al., 2002). C5b-9 complexes were not observed in soft drusen 549 (Mullins et al., 2014). 550 On whole-mount hydrated preparations of the choroid and BrM, (hard) drusen were 551 located to the intercapillary pillars of the choroid, suggesting a close relationship 552 between drusen formation and the capillary bed (Lengyel et al., 2004). This was 553 observed in earlier studies, but not systematically examined (Friedman et al., 1963). It 554 was suggested that drusen are a manifestation of (a) disturbed transport mechanism(s) 555 of substances across the capillary wall or BrM (Penfold et al., 2001). Whether this 556 indicates that drusen deposition is the result of slower clearance at the intercapillary 557 pillars or a manifestation of a disturbed transport mechanism of substances across the 558 capillary wall, or both, needs additional investigation. Of note, further pathological 559 compromise of the vascular bed and BrM leads eventually to the development of 560 subretinal neovascularization and wet AMD. 561 3.2.3. Contribution of blood proteins. 562 Penfold and coworkers suggested that breakdown of the normal choroidal vascular 563 function allows the movement of plasma proteins to the sub-RPE-BL space and this 564 leakiness is one of the cause of initiating the progression to AMD (Penfold et al., 2001). 565 Another study involved the analysis of age-related changes in various proteins and lipids 566 in the BrM using multiplexed Raman spectroscopy and found age dependent change in

heme signals (Beattie et al., 2010). However, there are no detailed and definitive studies how these plasma molecules end up in the sub-RPE-BL space. Involvement of fenestrations, breakdown of tight junctions, active vesicle transport (caveola) and receptor-mediated endocytosis (for macromolecules) have been suggested. Fenestrations are found predominantly on the endothelial vessel wall closest to the RPE (Bernstein and Hollenberg, 1965; Pino, 1985; Mancini et al., 1986). Rodent studies suggested that the number of fenestrae initially increases with age; but in advanced age and in AMD the number of fenestrae decreases (Burns and Hartz, 1992; McLeod et al., 2009). Transport through fenestrae is likely to be tightly regulated but it is not yet fully characterized (Pino and Essner, 1981; Essner and Gordon, 1983). Tight junctions of the choroidal capillaries show a tendency to become leaky with age, and lack transport regulation which may facilitate movement of plasma proteins from the choroid towards to the RPE (Nakanishi et al., 2016) (Aiello et al., 1998). Finally, vesicle- or receptormediated transport of proteins also exist in the choroid. (Smith et al., 1989). Taken together, transport of proteins at the choroid/BrM interface is complex and warrants further investigation. It has long been speculated that both blood plasma and incomplete digestion of photoreceptor outer segments contribute to the buildup of drusen material (Farkas et al., 1971a). It has also been suggested that drusen formation in the retina may be similar to plaque formation in arterial walls (Curcio et al., 2001), which, again, suggests that the contribution of blood proteins may be more important than previously thought (see section 8 on "drusen and plaques"). However, there is a paucity of information as to what extent proteins from the blood really contribute to drusen formation. It is thus plausible that some molecules exit the choroidal vessels into the extracellular space adjacent to the RPE, especially as the barriers in place to prevent such an event from

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happening, become compromised with age.

594 4. Selection of transcriptomic and proteomic datasets to determine the origin of drusen 595 proteins 596 4.1. Exclusion criteria and considerations. 597 One of the main goals of this study was to compare subretinal cellular transcriptomics 598 and proteomics as well as the blood proteome with proteins that are present in drusen. 599 To achieve this, we made use of a subset of studies from the literature as well as our own 600 data. Apart from the drusen protein studies, which date back to 2002, we only 601 considered here mRNA and protein studies published over the last 8 years; we did not 602 include retinal microRNA studies, non-coding RNA, metabolomics, imprinting studies 603 and data from (differences in) single-cell expression studies, simply because there are 604 relatively few confirmed and validated studies for the various types of retinal tissues 605 available yet. 606 Multiple excellent transcriptomics and proteomics studies have been published on 607 different layers of the retina/RPE/choroid complex, these are reviewed by a number of 608 authors recently (Skeie and Mahajan, 2014; Tian et al., 2015; Zhang et al., 2015a). 609 However, the studies currently available differ in many aspects, including study design, 610 retinal area and retinal cell type examined, sample source selection, sample handling, 611 sample numbers investigated, probe labeling methodology, microarray- or RNA 612 sequencing- methodology as well as the platform, quality and type of bioinformatics 613 programs used for analysis. It is not our goal here to describe and compare all the retinal 614 transcriptomic or proteomic data in the literature. Nonetheless, if one wants to compare 615 different sources (subretinal transcriptomics and proteomics, blood proteomics) and/or 616 outcomes (drusen proteins), similarity of the components and parameters of the 617 comparison(s) is obviously, highly desirable (Ahmad et al., 2018). 618 In the relevant transcriptomics literature, at least three phases can be observed: studies 619 before and after the introduction of the MIAME (Minimum Information About a 620 Microarray Experiment) quality guidelines studies (Brazma et al., 2001); studies before 621 and after the introduction of whole genome microarrays (at least 22000 genes (22 K or 622 more)) and studies before and after the introduction of RNA-Seg and GTex criteria. Over 623 time, a similar technological development has taken place in the proteomics field: from 624 2-D gels to high pressure liquid chromatography columns coupled and high throughput

625 mass-spectrometry-based studies (Geyer et al., 2016). In principle, the quality of large-626 scale transcriptomics and proteomics studies has continued to improve, and better and 627 more complete datasets may become available in time that may change some of the 628 interpretations described here. 629 There are several obvious differences between transcriptomics and proteomics studies. 630 In principle, transcriptomics techniques are highly sensitive and highly quantitative, but 631 as such, highly susceptible to RNA contamination or degradation. In addition, 632 transcriptome changes may not equate with changes on coded proteins and as such are 633 further away from biological function. 634 Proteomics studies, however, are usually less sensitive and quantitation can only be 635 achieved under certain circumstances, but proteomes per se are closer to function. 636 During disease progression, transcriptomics and proteomics profiles of a tissue can 637 change rapidly depending on disease stage. Also, a single tissue under study can be 638 affected by two or more consecutive disease stages at the same time. For example, in 639 AMD, new hard drusen continue to appear in the sub-RPE-BL space, while other drusen 640 in the same tissue already become confluent, and perhaps part of the same retina is 641 already prone to neovascularization. Consequently, for a disease like AMD, where the 642 RPE is subject to consecutive, insidious and overlapping disease stages, it is very difficult 643 to sift out useful and consistent healthy and disease stage specific expression profiles for 644 this cell layer. 645 Obviously, transcriptomics and proteomics studies cannot be translated one-to-one, due 646 to, for example, differences in RNA and protein synthesis and turnover rates. The sound 647 interpretation of both transcriptomics and proteomics is highly dependent on the use of 648 advanced bioinformatics and knowledge databases, which combine millions of data-649 points from human, mouse, and rat studies. Nonetheless, it is the investigator, with 650 knowledge of disease pathology, molecular biology and bioinformatics alike, who can 651 make the difference. 652 There are two goals with most transcriptomics (or proteomics) studies: One type of 653 study aims to find a complete molecular blueprint of the cells or tissues of interest; these 654 studies usually yield an enriched expression data set for the cell of interest. This type of 655 study usually includes both genes specifically expressed in the cell type of interest, but 656 also genes expressed in similar cell types. For example, the RPE is probably defined by a 657 few hundred RPE-specifically expressed genes, a few thousand neural cell-type

658 expressed genes, many expressed housekeeping genes for basic functions, as well as 659 many genes which are on "standby". The genes that are on "standby" have a very low 660 (leaky) expression if the cell in is a state of homeostasis. However, if the environment 661 changes, these very low expressed genes can rapidly be expressed to adapt the cell to a 662 changing environment. For example, the RPE shares most likely the RNA expression of a 663 large portion of its transcriptome: neural cell type genes, the household genes, and low-664 level expressed genes, with the other (neural) cell types in the retina (own 665 observations). Finally, there are many specific non-expressed genes in a certain cell-666 type. An example of expression studies which aim to find a molecular blueprint of the 667 cell is the uncurated RPE expression dataset, RPE-ET (Table 3), which contains 10% of 668 the biologically highest expressed genes in the RPE (Booij et al., 2009). 669 The other type of study aims to find only a maximum of genes specifically expressed in only the cells or tissue of interest. These few hundred genes, in the context of the more 670 671 generally expressed genes, give the cells of interest their specific cell type-associated 672 functionalities. An example is the dataset, RPE-ST (Bennis et al., 2015), which contains 673 170 RPE-specific expressed genes derived from previous RPE expression studies (Booij 674 et al., 2009; Booij et al., 2010b; Strunnikova et al., 2010) (STable 2). 675 4.2. Description of expression datasets used for drusenomics. 676 Apart from the 89 drusen protein data set, we used in this review 11 additional 677 subretinal and blood data-sets derived from previous transcriptomics and proteomics 678 studies; this is summarized in Table 3. We found that these transcriptomics and 679 proteomics databases complement each other and, together, give a more complete 680 overview of relevant expressed genes/proteins per tissue investigated. A common 681 feature of all high throughput studies is that they generate, by default, a small 682 percentage of misidentifications. This is due to cellular or molecular contaminations, or 683 mis-representation due to experimental sample handling. Therefore, individual gene 684 findings usually need to be confirmed by at least a second technique which focuses on 685 the analysis of single genes or proteins. 686 We used pure, enriched and curated cellular expression datasets. Pure datasets are 687 those without possible contaminations of other cell types while enriched datasets are 688 those datasets that have a certain degree of contamination of adjacent cell types. Finally, 689 curated datasets are those which are manually enriched either by bioinformatics or by

691 possible. The curation strategies employed are presented in Figure 3. 692 Most of the (non-curated) data were used for qualitative studies, have been published 693 and analyzed elsewhere, and are mentioned below for reference. For the quantitative 694 studies, we used curated datasets. The photoreceptors and choroidal transcriptome 695 datasets, cPR-ET and cChor/ET (Booij et al., 2010b) (GEO database accession number 696 GSE20191) have not been fully published before and therefore, their description will 697 receive a little more attention here. 698 699 First of all, we used (1) a combined data set for drusen proteins, curated by hand as 700 described above (Table 1). Furthermore, we used (2) a photoreceptor outer segment 701 proteomics dataset published by Kiel and coworkers (Kiel et al., 2011), which contains 702 proteins reflecting a multiscale signaling network associated with rhodopsin, the major protein component of rod photoreceptor outer segments. It was constructed by 703 704 combining relevant proteomics datasets, structural and functional literature mining and 705 bioinformatics approaches (Table 3; STable 3). Most likely, this database listing contains 706 some contamination from adjacent cell types, the RPE and choroid. Therefore, a curated 707 list was used for the quantitative studies: we subtracted the most highly expressed 708 sequences of the choroid (top 10% chor Booij; Chor-ET; and the uniquely expressed 709 sequences of the RPE (RPE-ST, Bennis)) from this database listing. The acronym used for 710 this dataset in this manuscript is PRos-EP (Photoreceptor outer segment-enriched 711 proteomics). The annotation of the curated version (c) of this dataset is cPRos-EP. 712 (3) The RPE-specific database with 170 entries was constructed by bioinformatic 713 curating and combining other (highly) enriched RPE gene expression databases (Booij et 714 al., 2010b; Strunnikova et al., 2010; Bennis et al., 2015). This database listing should be 715 viewed as a minimal number of RPE-specific expressed genes based on previous –omics 716 studies; the acronym used here is RPE-ST (Specific Transcriptomics) (Table 3; STable 2). 717 (4) The RPE secretome data from (Pao et al., 2018) that was published recently. RPE 718 cells were grown in vitro to confluency while adding various amounts of zinc to the 719 culture medium. Both the apically and basally secreted RPE proteomes were 720 determined. Here, we use the basal secretome proteomics listing which contains 276 721 entries. (Table 3; STable 1). Due to its nature, this dataset does not contain 722 contamination from other cell types but may contain contaminants from the culture

literature search to remove inevitable contaminations or irrelevant data as much as

- medium. In addition, its *in vitro* basis may not be fully representative of the *in vivo*
- situation, particularly in the disease state. The acronym for this database in this study is
- 725 RPE-IVS (*in vitro* secreted) (Table 3).
- 726 (5) The RPE/choroid proteomics dataset from Zhang and coworkers that contain
- 727 proteins extracted from RPE/choroid tissues of eyes from five individuals, fractionated
- and separated using SDS-PAGE and analyzed using mass spectrometry (Zhang et al.,
- 729 2016). In the RPE/choroid the authors identified 2755 non-redundant proteins. This
- dataset is rather large in components and is likely to contain entries from multiple cell-
- types (RPE, choroid, blood and possibly PR), and not only (RPE/choroid), given the
- inevitable contaminations of the PR sample with RPE and *vice versa*, and the
- contamination of the choroid with blood. The authors deposited their data to the
- 734 ProteomeXchange Consortium via the PRIDE partner repository with the dataset
- identifiers PXD001424 and PXD002194. The acronym for this database in this
- manuscript is RPE/chor-EP (<u>RPE/chor</u>oid-<u>e</u>nriched <u>proteomics</u>) Table 3.
- 737 (6) The blood proteome listing by Geyer and coworkers was produced by a new efficient
- 738 plasma proteome profiling pipeline (Geyer et al., 2016). Using a modified mass
- 739 spectrometry-based workflow they were able to identify and quantify at least 1000
- plasma proteins. Given the nature of the samples, it is unlikely to contain other retinal
- cells or proteins as contamination. The acronym for this database in this study is BL-SP1
- 742 (Blood plasma-specific proteomics; no 1) (Table 3; STable 4) (Geyer et al., 2016).
- 743 (7) The blood proteome dataset by Farrah and coworkers contains a non-redundant set
- of 1929 protein sequences from human plasma detected by tandem MS (Farrah et al.,
- 745 2011). The full data are available via PeptideAtlas, a large, international database of
- publicly accessible peptides identified in tandem MS experiments in a multitude of
- organisms. This is also a "pure" database listing. The original dataset contains
- endogenous chemicals, which we removed for our analyses. The acronym for this
- 749 database in this study is BL-SP2 (Blood plasma-specific proteomics, no 2); (Table 3).
- 750 (8) The BL-PHP blood proteome dataset consists of 262 HAP binding proteins from AMD
- patients and controls, as recently described (Arya et al., 2018). Plasma samples were
- taken from 23 individuals aged 65-90 with late stage AMD, each displaying drusen and
- 753 choroidal neovascularization in clinical images and attending the anti-VEGF injection
- 754 clinic at Moorfields Eye Hospital, London (STable 5).

- 755 (9) The atherosclerosis plaque proteomics dataset contains 3196 entries based on a
- 756 comprehensive review of the literature in this field (Bleijerveld et al., 2013). The
- acronym used in this study is AS-EP (Atherosclerosis-enriched proteomics) (Table 3).
- 758 The (large) dataset is available as supplementary file to the authors' publication.
- 759 (10-12) Transcriptomics datasets of the photoreceptor (acronym: PR-ET:
- 760 Photoreceptor; enriched transcriptomics), the choroid (acronym: Chor-ET: Choroid-
- 761 enriched transcriptomics.) (Table 3), and RPE (acronym: RPE-ET: RPE-enriched
- 762 <u>transcriptomics</u>) were produced using the same Agilent methodology and platform. For
- functional annotation and quantitative analyses, curated versions of these databases
- were constructed, named, respectively, cPR-ET (STable 6) and cChor-ET (STable 7). The
- 765 (c)RPE(-ET) database has been extensively published elsewhere (Booij et al., 2009; Booij
- 766 et al., 2010b).
- 767 4.3. Functional annotation photoreceptor (cPR-ET) and choroidal (cChor-ET) datasets.
- The PR-ET and Chor-ET datasets contain, respectively, the averaged top 10% highest
- 769 expressed genes in the photoreceptor and choroid. The isolation methods, study design
- and methodological issues for these datasets have been extensively discussed elsewhere
- (Booij et al., 2009; Booij et al., 2010b). These raw datasets were used for the qualitative
- studies in this manuscript. The experimental studies were performed in agreement with
- the declaration of Helsinki concerning the use of human material for research and
- followed both MIAME and GTex criteria (Brazma et al., 2001; Consortium, 2013).
- We curated both datasets PR-ET and Chor-ET according to scheme C in Figure 3. In
- order to obtain cell-specific datasets for photoreceptor and choroid, which are useful for
- both cell-specific functional annotation and for quantitative studies described elsewhere
- in this manuscript. Consequently, we removed from the PR-ET and Chor-ET datasets
- all expressed genes that overlap between them (either contaminations or truly shared
- gene expression). This resulted in two smaller curated datasets. Subsequently, we also
- 781 removed all potentially present RPE-expressed unique sequences (RPE-ST dataset) to
- generate the cPR-ET and cChor-ET datasets. Thus, the resulting cPR-ET and cChor-ET
- datasets contain less, but highly cell-specific entries compared to PR-ET and Chor-ET.
- Hence, we ended up with a highly photoreceptor-enriched gene expression dataset
- consisting of 745 genes (STable 6) and a highly enriched expression dataset for the
- 786 choroid of 848 entries (STable 7).

787 We ran an Ingenuity core analysis (www.ingenuity.com) on both cPR-ET and cChor-ET 788 datasets. This type of analysis typically yields data-driven functional annotations (i.e. it 789 produces biological motifs, canonical pathways and molecular networks enriched in the 790 dataset). The results of the cPR-ET analysis are presented in Table 4; PDF summary. We 791 found some very basic and very specific functional features related to established 792 photoreceptor function. The basic annotations included "cancer", "cellular function and 793 maintenance" as well as "tissue morphology". One could speculate that these relate to 794 the unique shape of the photoreceptor cell, and its unique ability to renew its 795 photoreceptor outer segments. More specific (highly ranked) annotations included 796 "photo transduction cascade", "visual system development and function" and 797 "neurological disease". These data-driven results clearly fit with reported specific 798 photoreceptor functionalities from the literature (Diamond, 2017; Musser and Arendt, 799 2017; Fain and Sampath, 2018). 800 The choroidal transcriptomics dataset cChor-ET was generated in a similar way to the 801 photoreceptor cPR-ET dataset described above. Obviously, the choroid is not a single 802 tissue, but consists of multiple cell types, including endothelial cells, fibroblast cells, 803 melanocytes, macrophages, and resident lymphocytes. The choroid is unavoidably 804 contaminated with blood cells and proteins. Nevertheless, after curation, we obtained 805 848 genes with a highly enriched choroidal expression in the cChor-ET dataset (STable 806 7). Following Ingenuity core analysis, the resulting functional picture of the choroid is, as 807 expected, completely different from that of the photoreceptors (Table 5; PDF summary). 808 We found that two of the top five biological motifs ("inflammatory response" and 809 "inflammatory disease") and three canonical pathways ("antigen presentation pathway", 810 "acute phase response signaling" and "complement system") are all involved with the 811 immune system. This confirms the crucial role of the choroid and blood in external 812 immune surveillance of the eye (Dick, 2017). The second highlight of this analysis was 813 the canonical pathway "atherosclerosis signaling" which again points to an important 814 resemblance between healthy or disease processes going on at the BrM (choroidal-RPE 815 interface) and the vessel walls (see also section 8). 816 Finally, both the canonical pathways and the highest ranked networks identified in this 817 cChor-ET analysis indicate tissue damage and injury. One possible explanation is that 818 this damage refers to early molecular complement attack already present or setting in, 819 which may be well before any morpholocial changes or damage may be visible.

Alternatively, although we used data from healthy post-mortem eyes, performed the
studies according to MIAME and GTex guidelines, 3' primer design which avoids
potential problems due to 5' directed degradation, as well as very stringent RNA quality
controls, the tissue damage and injury might still be due to post-mortem damage.
Detailed data relating to both cPR-ET and cChor-ET analyses are available on request.

826 5.1. Comparative study design considerations. 827 From the previous sections, it has become clear that a systematic investigation into the 828 origin of proteins in drusen is lacking in the literature. Authors suggest a variety of 829 protein sources, frequently on the basis of single observations. Systematic investigation 830 of this phenomenon is hampered by the heterogeneity of source samples, methodology, 831 and analysis. Large scale transcriptomics or proteomics studies frequently end up with a 832 rather abstract annotation analysis, allowing a certain error rate and lack of detail; small 833 scale studies frequently lack sufficient technical, methodological or biological replicates. 834 In a first attempt to investigate the origin of drusen-proteins systematically, we used the 835 presence of functional protein clusters identified in drusen (described in section 2). 836 Subsequently, we investigated whether expression of entities in these clusters also 837 (partly) occurs in the various non-curated expression database sets selected for this 838 study. An overview of this comparison is presented in Table 3. This comparison serves 839 two purposes: Firstly, the presence or absence of clusters in subcellular databases or 840 blood may give a qualitative indication of the origin of (the proteins in) the cluster and 841 secondly, it gives an indication of the completeness, quality, and contamination in each 842 of the databases listed. 843 We argued above (section 4.2.) in detail that all individual transcriptomic and 844 proteomics database lists used here (and those in the literature) are incomplete and, as 845 a rule, have a degree of RNA/protein/cellular contamination due to original mixed cell 846 sampling. How do we compare incomplete, contaminated datasets? 847 First of all, one should have some knowledge of the study-design and character of the 848 dataset under study, to understand why certain entries do, or do not, appear. The 849 characteristics of the databases used are described in section 4.2 above. As an example, 850 in the study of RPE/Chor-EP expression dataset (Table 3) we systematically identified 851 that a large number of functional cluster queries/entries are indeed present. However, 852 this is most likely due to the fact that the RPE/Chor-EP study contains proteins from 853 photoreceptor, RPE, choroid and blood. Thus, we decided to use this dataset as a positive 854 control (i.e., a dataset where almost all genes relevant to the study are 855 expressed/present). Similarly, we used either the unique sequences from the RPE-ST

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5. Drusenomics, part II: Qualitative analysis.

856 dataset, or the consequent absence of an entry in a set of similar dataset listings, as a 857 negative control. 858 In our qualitative comparison, the incompleteness and potential contaminations of the 859 various non-curated datasets (Table 3) may be largely overcome by considering similar 860 data from different studies at the same time. For example, a specific query may be 861 present in all photoreceptor studies, and, at the same time be absent from all blood 862 proteomics studies. 863 5.2. Where do proteins in drusen come from? A qualitative comparison. 864 We will now turn to the interpretation of the highlights of the comparative study results 865 presented in Table 3. In the first column (top to bottom) the functional gene clusters 866 from the molecular networks of drusen (Figure 2a-2d) are presented together with their 867 individual gene content (column 2) and functional annotation (column 3). On the top 868 rows (fourth column onward), subretinal transcriptomics and proteomics as well as 869 blood proteomics data(sets) from the literature are given. For full length names of the 870 abbreviated gene/proteins, see Table 1. 871 5.2.1. Network 1.1: The complement gene cluster. 872 The first functional cluster (Table 3; Figure 2a), in our analysis consists of the 873 complement end proteins: C7, C8 isoforms as part of the/and the Membrane Attack 874 (MAC) protein group in general as well as the multifunctional PRELP protein. Mutations 875 in PRELP cause myasthenic syndrome (Engel, 2018). Among other functions, PRELP is 876 involved in regulation of the complement cascade (Engel, 2018). As expected, we 877 observed that the (alternative pathway) complement gene transcripts/proteins are 878 absent from all photoreceptor and/or RPE transcription and proteomics datasets (PRos-879 EP, PR-ET, RPE-ET, RPE-ST, RPE-IVS). The only possible exception is the presence of 880 these proteins in the RPE/chor-EP dataset (positive control), where they most likely 881 originate from the choroid/blood component of the sample (please note again that the 882 choroid sample is inevitably contaminated with blood). In contrast, in 2 out of 3 blood-883 plasma datasets (BLP-SP1 and BLPHP) this cluster is present, except for PRELP. The 884 latter entry is apparently uniquely present in the Chor-ET listing (and in the positive 885 control RPE/Chor-EP) and is probably produced in the choroid. Interestingly, this leads

us to suggest that the systemic driven complement attack from the blood is locally

regulated by PRELP produced by choroidal cells (Happonen et al., 2012).

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389	5.2.2. Network 1.2: The collagen cluster.
390	The second functional cluster is that of the collagens and related molecules (Table 3;
391	Figure 2a). Here, the picture directly becomes, perhaps understandably, more
392	complicated. Collagen proteins are likely to be produced (RNA, protein) by the basal
393	secretion of the RPE and apical secretion of choroidal cells, and become part of the a-
394	cellular BrM (Booij et al., 2010a) and, at least theoretically, by the apical RPE and by the
395	PR for the interphotoreceptor matrix (IPM). Indeed, proteomics of apical RPE secretion
396	in vitro (Fort and Lampi, 2011) and functional annotation of the in vivo human RPE and
397	photoreceptor predicted secretomes (based on transcriptomics of the RPE and
398	photoreceptor cells; Bergen, unpublished), suggest that several specific collagen
399	proteins may (transiently) be present in the IPM, although their presence was never
900	detected by immunohistochemistry (yet).
901	During life, there is a constant turnover of ECM's, resulting in a mix of newly synthesized
902	and (partly) digested collagen fragments shuttling around the subretinal area.
903	Interestingly, the unique RPE-ST database (170 entries) does not contain any collagen
904	related entries, thereby confirming that, if the RPE produces collagen, none of these
905	collagens are made exclusively by the RPE (but also by adjacent tissues like the choroid).
906	Further to this, it is of interest to note that collagen (type $8A1$) is produced or present in
907	the PR-ET, RPE-ET, Chor-ET and the RPE/CHOR-ET datasets, which supports the
908	hypothesis that the (two collagenous layers of the) BrM, at least in part, are built from
909	both the RPE and choroid sides side (Booij et al., 2010a). The data in Table 3 further
910	suggest that the BrM proteins COL1A2, COLA1 and COLA2 are produced or are present
911	exclusively (or at least mainly) in the Chor/blood, and not in the PR and RPE datasets.
912	COL6A1 is secreted basally by the RPE <i>in vitro</i> (Pao et al., 2018) (Table 3), and is part of
913	the BrM (Booij et al., 2010a). Consequently, the protein may end up in the drusen
914	dataset either as a contamination, or as a remnant of the turnover of BrM components.
915	The proteins THBS4 and TNC occur only in one of the blood proteomics datasets (BL-
916	SP1). The presence of other entities (RBP3, EFEMP1, PLG, GPNMB, SEMA3B, TBHS4)
917	from this cluster in multiple PR/RPE and Chor/blood listings suggest that these
918	genes/proteins can be derived from different sources.

920	The third cluster of drusen proteins to be discussed are the crystallins (Crabb et al.,
921	2002; Nakata et al., 2005) (Figure 2a), frequently referred to as heat-shock proteins,
922	which act as chaperones to prevent or reduce protein degradation in stressed or aging
923	cells. Although they may have a more structural role, it is possible that the expression of
924	crystallins is increased only in those studies in which cells or tissues have been exposed
925	to a relatively large amount of stress. This would mean that further systematic and
926	methodological analysis of consistent (differences in) expression does not make sense. It
927	is remarkable however, that the CRYBB2 protein is present in the PRos-PT, RPE-IVS, and
928	the RPE/Chor-EP studies and in drusen. Consequently, this protein may originate from
929	the PR outer segments, processed, transported and secreted by the RPE and then
930	accumulates in drusen. To our knowledge, this is the only photoreceptor protein known
931	to possibly make it through phagocytosis and lysosomal processing in the RPE and end
932	up in drusen (Feeney-Burns et al., 1988; Hoppe et al., 2001).
933	5.2.4. Network 2.4: Genetic and developmental ophthalmic disorders.
934	The fourth functional gene cluster (Network 2, Figure 2b) can be considered as a
935	pathobiological cluster of developmental and ocular disease. The comparative analysis
936	(Table 3) shows that ATP5F1B, ACTB and annexin2 (ANXA2), are present in a number of
937	PR/RPE and Chor/blood expression datasets. These entries may thus be expressed in
938	multiple cell types or blood. ANXA2 is included here "within brackets", since it was
939	initially assigned to drusen using proteomics, but later the same authors stained for
940	ANXA2 in human donor eyes and concluded it was not present in drusen (Crabb et al.,
941	2002; Nakata et al., 2005). The CRYAB, ENO2 and SPTAN1 proteins cannot be clearly
942	assigned, but appear to be of a local cellular origin (PR, RPE, or Chor) and not from the
943	blood. The subcellular/systemic assignment of the FN1 and MYH9 entries are not clear.
944	The BFSBP1 and BFSP2 proteins neither occur in the subretinal datasets, the blood
945	proteomics lists, nor the positive control (RPE/Chor-EP; Table 3). According to the
946	literature, both are structural proteins that specifically form filaments in the
947	cytoskeleton of lens-cells (www.ingenuity.com). We therefore conclude that these are
948	very weakly expressed genes which express proteins that build up slowly and/or with a
949	long half-life. The only other explanation that could be offered is that they are
950	contaminations within the drusen dataset.

5.2.5. Network 3.5: Injury, inflammation and dermatological disease.

952 The fifth functional drusen cluster (Network 3; Figure 2c) is related to interacting genes 953 and proteins involved in injury, inflammation and dermatological disease. A substantial 954 number of entries of this group seem to have both a cellular as well as a systemic 955 presence or origin since they are present in at least two database listings from PR/RPE 956 and Chor/blood category. These include ANXA1, ANXA5, CKB, GAPDH, PRDX1 and 957 S100A8. 958 The SERPINA3 and ALDH1A1 proteins appear only in at least two of the RPE-Chor-EP, 959 Chor-ET, BL-SP1, BL-SP2 and the BL-PHP datasets, but not in the PR/RPE dataset. Thus, 960 both proteins appear to come from the systemic side of drusen. FRZB (SFRP3) 961 (www.genecard.org) is present in 3 PR/RPE listings (Table 3), including the RPE-specific 962 listing (RPE-ST), and in only one choroid-enriched list (Chor-ET). We tentatively assign 963 this drusen protein primarily to the RPE, and as a contamination in the (PR and/or) Chor 964 database listings. S100A7 is only once present in the PRos-EP proteomics dataset. 965 Assignment of S100A9, TYRP1, LAMB2, APOE, and FrzB or LUM to a single source cannot 966 be done on the basis of this comparison. 967 5.2.6. Network 4.6: Cell to cell signaling; systemic involvement. 968 As can be expected from the functional annotation "cell to cell signaling; systemic 969 involvement", almost all of the entries of this category of drusen proteins, appear in the 970 Chor/blood datasets (Table 3). The exceptions are clusterin (CLU), ANXA6 and HRG. 971 From the literature, we know that CLU is a ubiquitously expressed gene that is 972 expressed in all cell types (Wilson and Zoubeidi, 2017). It is therefore not surprising that 973 it features in both the PR/RPE as well as the Chor/blood listings. The final assignment of 974 ANXA6 and HRG, on the basis of this comparison is not clear. The role of the systemically 975 derived HRG drusen protein is discussed in detail below (section 7). Of particular 976 interest is the expression of CFH, given its central regulatory role in the complement 977 attack on (chemically modified) drusen components. There is compelling evidence in the 978 literature that CFH is present in the blood, the neural retina and that it is also expressed 979 by the RPE (Li et al., 2014; Mullins et al., 2014; Whitmore et al., 2014; Chirco et al., 2016; 980 Chirco and Potempa, 2018; Toomey et al., 2018). The presence of CFH protein in the 981 blood corresponds with the data and proteomics listings of blood in Table 3. What is not 982 entirely clear is why CFH does not pop up in the RPE listings. This can perhaps be 983 explained as follows: The enriched RPE-ET transcriptomics list contains only the highest

984 10% expressed genes in the RPE. Apparently, CFH is somewhat lower expressed and so 985 does not belong to this group (Warwick et al., 2014). Also, the RPE-ST specific listing 986 only contains 170 entries uniquely expressed by the RPE; whilst CFH is produced in 987 other cells or blood as well. Finally, CFH does not occur in the RPE-IVS basal secretion 988 proteomics listing, which is not entirely unexpected as recent evidence suggests that 989 CFH is secreted apically, not basically by the RPE (Kim et al., 2009; Pao et al., 2018). 990 5.2.7: Conclusion. 991 On the basis of our qualitative comparison, we suggest that a number of (functional 992 clusters of) drusen proteins come from the blood, while others come from a subretinal 993 cellular compartment. These results are in line with the findings in the literature. 994 However, it is not clear yet how many of the drusen proteins come from each particular 995 compartment. The latter may be estimated by a more quantitative analysis, which is the 996 subject of the next section.

6. Drusenomics, part III: A quantitative approach

6.1. Quantitative analysis and curation of datasets.

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1001 In this chapter, we quantitatively compare drusen proteins with transcripts and proteins 1002 from adjacent retinal compartments (photoreceptor, RPE, choroid) and blood. As 1003 described in section 2 above, the drusen protein list was compiled manually according 1004 to the curation strategy presented in Figure 3, scheme A. Similar to the qualitative 1005 studies, the quantitative analysis of the origin of drusen proteins is also hampered by 1006 two problems: (a) most large scale cellular transcriptomics and proteomics datasets 1007 contain some contamination (both RNA and/or protein) from adjacent cells or tissues, 1008 and (b) most of the datasets are incomplete due to differences in the study design and 1009 methodology used in contributing studies. In other words, we need to use a quantitative 1010 comparison strategy that maximizes the signal (number of entries to be compared) and 1011 minimizes the noise (number of contaminations in datasets). We overcame the 1012 incompleteness of various datasets by pooling the entries from various similar (cell-type 1013 specific) studies, to get a more complete numerical picture (Figure 3, scheme 3A). 1014 With regard to possible contaminations, we used two types of datasets. The first category includes datasets that, by definition or by previous curation in the literature, 1015 1016 contain cell-specific expressed entries only, such as the RPE-ST, RPE-IVS, BL-SP1 and the 1017 BL-SP2 datasets (section 4.2 and/or Table 3).. The curation of datasets PR-ET and Chor-1018 ET into cPR-ET (STable 6) and cChor-ET (STable 7) was already described above 1019 (section 4.2). The other category datasets used (PRos-EP, PR-ET, Chor-ET) were newly 1020 curated, as presented in Figure 3 and 4, in such a way that they, after curation, also only 1021 contained cell-type specific entries 1022 The PRos-EP dataset was curated according to the curation strategy presented in Figure 1023 3, scheme 3C: We removed from the PRos-EP dataset (in principle containing 1024 photoreceptor outer segment expressed genes only) all choroidal highly expressed 1025 genes (from Chor-ET) as well as potentially present uniquely RPE expressed entries 1026 (from RPE-ST) resulting in the curated cPRos-EP dataset (STable 6). The removed 1027 choroidal and RPE entries were (potentially) present in the PRos-EP dataset due to truly overlapping gene expression between these different cell types and/or due to 1028 1029 contaminations in the original cell sample. Of note, cPRos-EP does still contain entries 1030 from both photoreceptor and RPE since contamination between these two is inevitable.

1031	Together, the database listings cPRos-EP, cPR-ET, RPE-ST and RPE-IVS form the neural
1032	side of drusen database listings (Figure 4). Similarly, the cChor-ET, the BL-SP1 and the
1033	BL-SP2 constitute the systemic side of proteins found in the drusen dataset. In summary,
1034	we ended up with three large datasets suitable for further robust analysis: The drusen
1035	protein dataset, the "neural source of drusen" database, and the 'systemic source" of
1036	drusen dataset (Figure 4).
1037	Next, we compared the entries present in the "neural source listing" and in "the systemic
1038	source listing" with the proteins present in drusen. The result of this analysis is
1039	summarized in the Venn-diagram in Figure 5. The comparison revealed that 10 proteins
1040	appeared to be uniquely derived from the neural side (Table 6) and 37 proteins are
1041	derived from the systemic drusen side (STable 8). In addition, there were 23 proteins
1042	that come (potentially) from both the neural and systemic side (STable 9). For 19 drusen
1043	proteins (out of the 89), the origin remained unclear as they were neither present in the
1044	"neural source" nor in the "systemic source" expression datasets (STable 10).
1045	6.1.1. Ten out of 89 drusen proteins originate uniquely from the PR/RPE.
1046	Our analysis yielded 10 drusen proteins that originate from the PR or RPE (Figure 5;
1047	Table 6). They are both uniquely present in the drusen proteomics dataset and the
1048	neural source of drusen database listing. We traced these proteins back to their original
1049	source(s), and we observed that three of them (FRZB, RDH5 and RGR) originally came
1050	from the unique entries in the RPE-ST dataset, five came from the RPE-IVS dataset
1051	(CRYBA1, CRYBA4, CRYBB2, ENO2 and TUBB3), and the remainder from the cPRos; cPR-
1052	ET datasets. Taken together, 8 out of the 89 drusen proteins originated uniquely from
1053	the RPE, while 2 came from the curated PR/RPE database listings (cPROS; CPR-ET)
1054	(Table 3). Finally, we also reviewed the psychochemical properties and molecular
1055	weight (Mw) of these 10 proteins (Table 6). We do not know the HAP binding properties
1056	of these proteins, but they do not occur in the BH-PLP HAP-binders' dataset (STable 5).
1057	In conclusion,, we did not observe any common signatures of these proteins that would
1058	explain why they in particular are trapped in the sub-RPE-BL space (whilst other
1059	proteins are not).
1060	6.1.2. Twenty-three of 89 drusen proteins originate from both neural and systemic
1061	SOURCES.

1062 From our analysis, 23 drusen proteins were present in both the "neural source" as the 1063 "systemic source" datasets (Figure 5; STable 9). From these, only 1 protein (S100A9) 1064 falls in the curated PR/RPE category (Figure 4). Additional groups of two and twenty 1065 proteins come from the unique RPE RPE-ST and the RPE-IVS datasets, respectively. At 1066 the same time, all 23 of these proteins are also present in the blood. Remarkably, in this 1067 shared category the majority of proteins are either secreted basally by the RPE or 1068 present in a soluble form in the blood plasma. We hypothesize that the proteins in this 1069 category enter the sub-RPE-BL from both sides, where they "meet, greet and stick", i.e. 1070 form aggregates that cannot be cleared and therefore contribute to drusenogenesis. 1071 Functional and pathobiological annotation of (combinations of) these proteins can be 1072 found in STable 9a. 1073 6.1.3. Thirty-seven out of 89 drusen proteins originate from the choroid/blood. 1074 We found that 37 out of 89 drusen proteins uniquely originate from choroid or blood 1075 datasets (STable 8). From these, 31 proteins came from the plasma-proteomics datasets 1076 (BL1-SP1 and BL-SP2). The remaining six entries (ANXA6, FLBN5, HLA-DRA, MFAP4, 1077 PRELP, SEMA3B) are present in cChor-ET database listing and thus originate from either 1078 the choroid or blood. Functional and pathobiological annotation of (combinations of) 1079 these proteins can be found in STable 8a. 1080 In summary, we again observed a large proportion of drusen proteins are most likely 1081 originating from the blood. If we take this unique category (31 proteins from plasma) 1082 and the shared contribution of plasma (23 proteins) from the previous paragraph into 1083 account, we can conclude that as many as 54 out of 89 drusen proteins (>60 %) are (co-) 1084 derived from blood plasma. 1085 6.2. Nineteen drusen proteins out of 89 were not assigned. 1086 We can, in the end, still not determine the possible origin of 19 out of 89 drusen proteins 1087 (STable 10) Why is this not possible? Do these proteins have a number of characteristics 1088 in common that prevents us to determine their origin? To attempt to answer these 1089 questions we need to take a closer look at these remaining drusen proteins. The 1090 functional annotation of these proteins is presented in STable 10a and they can be 1091 divided in five groups: (1) a gamma-crystallin group; (2) a histone cluster group; (3) 1092 (remnants from) BrM turnover group (4) a beaded filament group; (5) a rest group 1093 containing a variety of proteins that do not belong to a specific functional group.

1094 In group one, we observed several gamma-crystallin-isoforms in drusen, which have not 1095 been assigned to a specific source (as yet). Crystallins are commonly found in the lens 1096 but are also present in soluble form in the retina (Jones et al., 1999) and probably act as 1097 chaperone proteins after (oxidative) stress. Indeed, in the mouse retina, crystallin 1098 expression has a binary nature in which either they are highly upregulated, or their 1099 expression is extremely low (Templeton et al., 2013). Gamma-crystallins may have a 1100 neuroprotective role (Thanos et al., 2014). At least one specific type of crystalline 1101 (alphaB type) is known to be secreted by the RPE through micro-vesicle release (Kannan 1102 et al., 2016). In conclusion, the source assignment of gamma-crystallin isoforms in 1103 drusen, in our comparison, may be hampered by this binary expression. More 1104 specifically, it will simply be absent from a number of subretinal expression datasets 1105 and, as such, too little evidence exists to make a definite assignment. 1106 Next, we found a group consisting of HIST1H1E, HIST1H2BJ, HIST1H2BL and 1107 HIST2H2BE. Histones are highly basic proteins that have an essential role in the 1108 maintenance of nuclear DNA structure and gene transcription. HIST1H1E is a 219-amino 1109 acid protein that binds to the linker DNA stretch between nucleosomes, while HIST2, 1110 together with HIST3 and HIST 4, are part of the nucleosome core (Tessarz and 1111 Kouzarides, 2014). Damaged or dying cells (potentially RPE or endothelial cells of the 1112 choroid) can release cellular as well as nuclear fragments that may contain histones. 1113 Alternatively, high concentrations of serum histones have been detected in several 1114 human diseases (Yang et al., 2015). These extracellular histones may get trapped in BrM 1115 and drusen. Interestingly, extracellular histones trigger activation of multiple signaling 1116 pathways related to cell death, growth and inflammation and may play a role in auto-1117 immunity, aging and disease (Allam et al., 2014; Kalbitz et al., 2015; Zhang et al., 2015b). 1118 Why these specific histone proteins (and not others) are trapped in drusen and cannot 1119 be assigned to a source remains to be elucidated. 1120 The third drusen protein group with an as yet unassigned source contains elastin (ELN), 1121 collagen 8A1 (COL8A1), biglycan (BGN) and tissue inhibitor of metalloproteinase 3 1122 (TIMP-3) proteins. These proteins may come from an as yet little considered drusen 1123 protein source: the BrM and its (turnover) components (Booij et al., 2010a; Curcio and 1124 Johnson, 2012). TIMP-3 is expressed in the RPE (Ruiz et al., 1996) and is crucial for the 1125 maintenance of BrM. Mutations in TIMPO-3 caused Sorsby Fundus dystrophy, a

1126 monogenic disease that resembles the phenotype of AMD (Weber et al., 1994). Indeed, 1127 as described previously, BrM is dynamic in nature, not only in a physiological sense, but 1128 its composition and properties vary with age. Proteins involved in BrM and its turnover 1129 may be absent from (some of) our subretinal transcriptomics and proteomics datasets, if 1130 the relative expression levels of such entries are low. The middle layer of the BrM 1131 consists of elastin so it is conceivable that the RPE and/or choroidal cells make this 1132 protein. Within the BrM, elastin turnover might be relatively low, thus little "new" 1133 elastin is needed. While elastin protein fragments (tropo-elastin) might be present in 1134 some drusen as a remnant from BrM-turnover, there is little evidence in the literature 1135 that they accumulate in drusen. 1136 Fourth, two members of the beaded filament structural protein family, BFSP1 and 1137 BFSP2, remain unassigned. Similar to crystallins, these proteins were initially 1138 discovered as lens fiber proteins. To our knowledge, it is not clear whether they also 1139 play a role in retina/RPE or maybe even the BrM. How these proteins end up in drusen 1140 and their origin remains unclear. 1141 The fifth, yet unassigned group contains a number of, apparently, unrelated proteins, 1142 including retinol binding protein 3 (RBP3), tyrosinase protein-like 1 (TYRP1), spectrin 1143 alpha, non-erythrocytic 1 (SPTAN1), disco interacting protein homologue (DIP2C), 1144 forkhead-associated phosphor peptide (FHAD1), and scavenger receptor class B 1145 member 2 (SCARB2). 1146 The RBP3 gene is transcribed in the PR and its protein is located in the interphotoreceptor matrix (IPM). It binds to retinoids which are shuttled from the PR to 1147 1148 the RPE, and vice versa (Gonzalez-Fernandez et al., 1993). The TYRP1 gene belongs to 1149 the tyrosinase family transcribed in the RPE and encodes an enzyme in the melanin 1150 biosynthetic pathway (Lai et al., 2018). Mutations in this gene are one of the causes of 1151 albinism (Kamaraj and Purohit, 2014; Kruijt et al., 2018). Both RBP3 and TYRP1 genes 1152 may be absent from our datasets given their relatively low, transient or binary 1153 expression in the relevant tissues. 1154 Finally, FHAD1 is a small protein that recognizes phosphorylated epitopes on a wide 1155 range of proteins as part of an evolutionarily ancient mechanism enabling assembly of 1156 protein complexes (Durocher and Jackson, 2002). The expression of FHAD1 is very low 1157 in many tissues including the retina but is high in the testis and lungs. Since the

1158 expression is very low, the transcript and protein production or presence may go 1159 undetected in the subretinal and blood transcriptomics and proteomics studies we used 1160 in this study. However, once the FHAD1 protein has accumulated, as apparently in 1161 drusen, it may be (more) detectable there. DIPC2 is a ubiquitously expressed protein 1162 that shares homology with a *Drosophila* protein that interacts with the transcription factor disco (www.ncbi.nlm.nih.gov). It is possible that the expression of this type of 1163 1164 protein is transient or binary; and it may go undetected in our retinal compartment and 1165 blood transcriptomics and proteomics studies for that reason. 1166 Finally, the last two proteins, SPTAN1 and SCARB2 may have related functionalities. The 1167 SPTAN1 protein is a part of the cytoskeletal spectrin protein family that is involved in 1168 stabilizing membranes of both cell and organelles (Tohyama et al., 2015). It is highly 1169 expressed in the brain, and still expressed to a significant level in multiple other tissues. 1170 SCARB2 is a ubiquitously expressed gene that encodes a lysosomal type III plasma 1171 membrane glycoprotein (Gonzalez et al., 2014). Given the involvement of this type of 1172 proteins in the lysosomal digestion of cellular material, it is tempting to speculate that 1173 these proteins come from (transiently present in high numbers) lysosomal membrane 1174 fragments basally secreted by the RPE. 1175 6.3. Blood proteins are an important source of drusen proteins. If we summarize the combined data from our literature search and our qualitative and 1176 1177 quantitative analyses, we conclude that blood proteins are an important protein source 1178 for drusen development. Further studies are needed to confirm and enhance our data, 1179 especially on the single-protein level. Given the apparent contribution of blood to the 1180 formation of drusen, the next chapters will discuss the role of hydroxyapatite as a 1181 retainer of blood proteins during drusen formation, and the similarities that exist 1182 between drusen and atherosclerotic plaques, which occur exclusively in the vasculature.

7. Drusen and hydroxyapatite

1184	Our analyses provide strong evidence that proteins in drusen come from multiple
1185	sources. The next logical step was to consider how proteins arrive and how they are
1186	retained in the sub-RPE-BL space. One possibility is that proteins may bind to
1187	constituents of the BrM (Tabas et al., 2007). Another is the formation of large oligomers
1188	in the sub-RPE-BL space in the presence of the high concentration of trace metals
1189	(Lengyel et al., 2007; Nan et al., 2013; Flinn et al., 2014). In addition, it was recently
1190	hypothesized that proteins might be retained in the sub-RPE-BL space due to their
1191	binding to hydroxyapatite spherules recently identified in human drusen (Thompson et
1192	al., 2015) (Figure 6). Since this hypothesis is relatively new, it is described in more detail
1193	below.
1194	Using confocal microscopy and hydroxyapatite (HAP)-specific fluorescent dyes, small
1195	hollow spherical structures ranging from 0.5 μm to 20 μm in diameter were identified
1196	within sub-RPE-BL deposits in retinal tissue sections of human cadaveric eyes
1197	(Thompson et al., 2015). The HAP spherules were present in all deposits examined
1198	(Thompson et al., 2015). Protein constituents of drusen, such as amyloid-beta,
1199	vitronectin and complement factor H, were localized to the surface of the HAP spherules,
1200	either individually or in combination (Thompson et al., 2015). Although not all
1201	investigated drusen proteins appeared to bind to the surface of HAP (Thompson et al.,
1202	2015), this finding proved that the retention of proteins can, at least partly, occur
1203	through this protein-HAP interaction. These results also suggested that the binding of
1204	proteins to HAP spherules is a wide ranging, though selective, process and thus
1205	understanding which proteins can bind to HAP might be important. The plasma protein-
1206	binding capacity and selectivity of HAP was recently examined using a quantitative
1207	proteomic approach called Sequential Window Acquisition of all theoretical fragment-
1208	ion spectra-Mass Spectrometry (SWATH-MS) (Arya et al., 2018). Using this approach,
1209	242 proteins with the propensity to binding HAP were identified and quantified (Table
1210	3; STable 5) (Arya et al., 2018). Taking advantage of the quantitative nature of the
1211	analysis the binding of samples from participants with wild type and the AMD associated
1212	high risk CFH variant, T1277C were compared. Quantitative differences in the
1213	abundance of at least 34 proteins were identified, suggesting that the genetic
1214	background is likely to affect the protein composition of drusen "simply" due to the

1215 availability of proteins in the blood. This approach also highlighted that there are 1216 proteins, whose presence and potential role in sub-RPE deposit formation and in AMD 1217 had not previously been explored. One such example is the pregnancy zone protein 1218 (Arya et al., 2018), a plasma protein whose levels are known to increase in pregnancy 1219 and some disease states such as AD (Nijholt et al., 2015). 1220 It appears therefore that while drusen deposition is a hallmark of AMD, HAP deposition 1221 is a hallmark of drusen formation. The study by Thompson and coworkers was not the 1222 first to identify calcified components of drusen (Thompson et al., 2015). Spherical 1223 particles of similar size were previously identified within drusen and BLinD (Green and 1224 Key, 1977), and electron microscopy (Ulshafer et al., 1987; van der Schaft et al., 1992; 1225 van der Schaft et al., 1993; Thompson et al., 2015). Particles size observed in these 1226 studies ranged from 0.5 µm to 10 µm in diameter and contained calcium and phosphate 1227 as determined by elemental analysis (Ulshafer et al., 1987). More recent studies using 1228 von Kossa staining, a silver enhancement technique that identifies phosphates salts also 1229 indicated that calcium phosphates were present within deposits in the sub-RPE-BL 1230 space (Suzuki et al., 2015). 1231 The precipitation of calcium phosphate from an aqueous solution is a complex process (Kani et al., 1983; Tas, 2000; Jang et al., 2014). At neutral pH, HAP is considered the most 1232 1233 thermodynamically stable form of calcium phosphate. In fact, it is possible that HAP is 1234 stable enough that once the lipid or protein components of the drusen regress (Sallo et 1235 al., 2009; Toy et al., 2013; Novais et al., 2015), HAP still remains and continues to 1236 interact with its environment. This may suggest that HAP interactions, not only with the 1237 BrM and the RPE but also with the remanence of photoreceptor cells or other parts of 1238 the neurosensory retina (Bird et al., 2014) may require further investigation. 1239 Figure 7 summarizes the model of HAP associated deposit initiation. Under normal 1240 1241 circumstances, there is a physiological exchange of material between the RPE and the 1242 choroidal circulation (Fig.7A and A'), and this includes the exchange of lipid particles 1243 (Curcio et al., 2011). With age and disease lipid particles start accumulating in the sub-1244 RPE space including the BrM (Fig.7B and B') (Curcio et al., 2011). In the presence of lipid 1245 droplets and homeostatic changes in calcium and phosphate availability in the sub-RPE-1246 BL space, HAP can precipitate on the surface of the lipid droplets (Fig.7C and C'). Then, 1247 on the surface of the HAP spherules drusen proteins can accumulate (Fig.7D and D') via

1248 directly interacting with HAP (Arya et al., 2018). Based on fluorescence labeling of HAP 1249 in human eyes, it is appeared that HAP spherules can exist without drusen (Fig.7C'), but 1250 drusen have not been seen without HAP spherules (Fig.7D and D') (when specifically 1251 looked for) thus far (Thompson et al., 2015). Based on these observations it was 1252 proposed that HAP deposition is a seeding point for drusen formation (Thompson et al., 1253 2015). 1254 The next obvious question to ask is where the HAP spherules are originating from? 1255 Could they be blood or RPE derived? Do they exist as spherules only in the sub-RPE-BL 1256 space or is the material present in the surrounding tissues? Spherules have not been 1257 detected in any of the cellular or intercellular spaces although the calcium phosphate 1258 crystals had been showed in mitochondria (Carafoli, 2010). Therefore, it appears that 1259 HAP spherules are deposited in the retina exclusively in the sub-RPE-BL space. In fact, it had been shown that HAP deposition can occur in primary RPE cell models which 1260 1261 showed that HAP deposition can be initiated by the RPE alone, although contribution of 1262 the culture medium cannot be ruled out (Pilgrim et al., 2017). Whether spherical 1263 structures can develop in a cell culture system that are co-cultured with endothelial cells 1264 and/or fed with photoreceptor outer segments will need to be investigated. 1265 HAP mineralization in the retina clearly differs from classical mineralization in bone, but 1266 it may, or may not, share some key features with general soft tissue/elastin calcification 1267 (Figure 8). Obviously, the retina lacks extracellular matrix forming osteoblasts. Also, no 1268 relationship has been found between spherule mineralization and general HAP 1269 deposition on elastin and/or collagen. However, systemic driven HAP deposition can 1270 take place in the BrM, as reported before (Gorgels et al., 2012). Indeed, a systemic lack of 1271 inorganic PPi in the blood (Jansen et al., 2013) may be involved in local HAP deposition 1272 in BrM, facilitated by local conditions, such as oxidative stress (Mungrue et al., 2011). 1273 Interestingly, investigation of the ultrastructure and composition of vascular micro-1274 calcifications associated with uremia showed the presence of spherical particles in the 1275 media of the kidney, with internal structures comparable to those observed in the 1276 human eye (Schlieper et al., 2010). Similarly, the loose stroma of the choroid plexus of 1277 the aging or Alzheimer's disease brain contain psammoma bodies, which are entities 1278 with distinct HAP cores and multiple concentric rings or swirls of collagen wrapped 1279 around it (Alcolado et al., 1986). More recently, similarly structured spherules were also

1280	identified within patients with osteoporosis and in cardiovascular disease (Bertazzo et
1281	al., 2013; Shah et al., 2017). Thus, comparable mineralization mechanisms in a variety of
1282	non-osseous tissues appear to be associated with a number of different disease
1283	conditions.
1284	Alternatively, transcriptomic data suggests that part of the elements of the physiological
1285	mineralization process are (also) present in the RPE cells (Booij et al., 2009), or at the
1286	RPE/choroid interface (Whitmore et al., 2014). This evidence suggests that the
1287	molecular machinery required for general physiological mineralization (depicted in
1288	Figure 8) could (also in part) be assembled in the outer retina. Given that the bulk of the
1289	calcium is extracellular, while phosphate is mainly localized intracellularly, the
1290	conditions that allow mineralization to happen could be present locally, in the sub-RPE-
1291	BL space. This concept is novel and has not been investigated previously but may lead to
1292	HAP-based treatment strategies and/or new early detection mechanisms.

1293	8. Drusen and plaques: age-related macular degeneration and atherosclerosis
1294	
1295	The finding that a substantial number of drusen proteins are blood-borne prompted us
1296	to re-summarize a possible relationship between initiation and propagation of drusen
1297	and atherosclerotic plaques. A possible link between these two diseases was previously
1298	suggested based on (controversial) epidemiological evidence, the involvement of similar
1299	lipoproteins in the formation of extracellular deposits in AMD and atherosclerosis and
1300	structural commonalities between the vessel wall and Bruch's membrane (Curcio et al.,
1301	2001; Sivaprasad et al., 2005).
1302	AMD is a disease starting with (multiple macular) drusen formation. Drusen consist of
1303	(oxidatively-modified) lipids and proteins as well as minerals (Sarks et al., 1988; Green
1304	and Enger, 1993; Curcio and Millican, 1999; Crabb, 2014; Flinn et al., 2014; Handa et al.,
1305	2017; Pilgrim et al., 2017; Spaide et al., 2018). Drusen constituents most likely invoke a
1306	complement attack and sustain a continuous low-grade inflammation, which leads to
1307	serious events such as RPE cell loss, neovascularization and ultimately, central vision
1308	loss (Bird et al., 1995; de Jong, 2006).
1309	Atherosclerosis is a disease associated with the build-up of plaques also composed of
1310	(oxidatively-modified) lipids, proteins as well as minerals in the vessel wall of arteries
1311	that, via complement attack and low-grade inflammation, can lead to serious events
1312	including heart attack, stroke or aneurysm (Simmons et al., 2016).
1313	There are a number of clear differences between AMD and atherosclerosis, such as
1314	location of the deposition, the local metabolic physiology (Stefansson et al., 2011),
1315	involvement of other (different) genes or molecules and obviously, aspects of the
1316	pathological consequences of deposition build up (Hageman et al., 2001; Hopkins, 2013).
1317	For example, other than in plaques in atherosclerosis, deposit (drusen) formation in
1318	AMD most likely blocks the exchange of biomolecules between the retina and the
1319	choroid. It is thought that this interferes with the "nourishment" of the sensory cells in
1320	the retina causing them to die. Over time the cells cannot be replaced leading to a loss of
1321	vision, typically within the macula, which progressively deteriorates over time
1322	contributing to the AMD pathology (Bhutto and Lutty, 2012).
1323	Nonetheless, a relatively large number of commonalities have been found between both
1324	drusen and atherosclerotic plaque formation and their associated diseases. Available

1325 evidence comes from clinical, epidemiological, genetic, histological and pathobiological 1326 investigations (see below). 1327 8.1. Clinical and epidemiological studies. 1328 Verhoeff and Grosmann were the first to suggest a relationship between vascular 1329 disease and AMD (Verhoeff and Grossman, 1937). Except for a few reports (Gass, 1967; 1330 Kornzweig, 1977), this observation was largely ignored for over forty years when 1331 Maltzman and Hyman (Maltzman et al., 1979; Hyman et al., 1983) pioneered a plethora 1332 of subsequent epidemiological studies on the subject (Vidaurri et al., 1984; Vingerling et 1333 al., 1995; Snow and Seddon, 1999). Some studies between atherosclerosis (and similar 1334 diseases) and AMD showed positive associations (The Eye Disease Case-Control Study, 1335 1992; Klein et al., 1993), while others did not (Hyman et al., 1983). These controversial 1336 results, especially in the early investigations, were partly due to differences in description of the clinical phenotype, use of different end phenotypes, study design, 1337 1338 population size, lack of suitable replication populations, and insufficient knowledge of 1339 possible confounders. Nonetheless, this issue has not been resolved up until today. 1340 A wide range of epidemiological studies have also suggested that there are certain risk 1341 factors which are common to both AMD and atherosclerosis (-associated cardiovascular 1342 disease). These most consistently include environmental factors, such as age and 1343 tobacco smoking (Woodell and Rohrer, 2014). These studies indicate that, 1344 mechanistically, oxidative stress and potentially lipid metabolism may play an important 1345 role in both disorders (Serban and Dragan, 2014; Gehlbach et al., 2016; George et al., 1346 2018; van Leeuwen et al., 2018; Wilson et al., 2018). 1347 8.2. Histological and pathobiological similarities. 1348 Histological and pathobiological similarities between drusen and atherosclerotic 1349 plaques and their associated diseases include similarities of lipid and mineral content 1350 and structural similarities between the BrM and the vascular wall, endothelial cell 1351 dysfunction, and proteoglycan turnover. Curcio and coworkers proposed, for the first 1352 time, a relationship between drusen and atherosclerotic plaques since both contain 1353 similar neutral lipids and both accumulate cholesterol esters (Curcio et al., 2001). This 1354 finding was confirmed by others (Chung et al., 2005; Wang et al., 2010). These and 1355 subsequent studies made clear that diseases related to this type of accumulations may 1356 be mediated by genetic variation, oxidative stress and inflammation. The accumulation

1357 of lipids in drusen and atherosclerosis has recently been reviewed elsewhere in detail (Pikuleva and Curcio, 2014; van Leeuwen et al., 2018; Xu et al., 2018). 1358 1359 Sivaprasad and coworkers observed that the BrM and the vascular intima share a 1360 number of common structural modalities, and age-related changes (Sivaprasad et al., 1361 2005). Indeed, similar to the vessel wall, and given the presence of local fenestrated 1362 choroidal capillaries, BrM acts a collagen and elastin rich physical barrier for the blood. 1363 Both the BrM and vascular intima thicken through accumulation of extracellular lipids 1364 and other debris and become less flexible with age (Chung et al., 2005; Curcio and 1365 Johnson, 2012). 1366 Another important feature of the ECM of both the vessel wall and BrM are the presence 1367 and turnover of a variety of proteoglycans. In BrM, the ratio between several proteoglycan types, most notably heparan sulfate and chondroitin sulfate, changes 1368 1369 dramatically during aging and the development of AMD (Barzegar-befroei et al., 2012). 1370 In atherosclerosis and AMD, (oxidatively) modified proteoglycans may bind and retain 1371 specific apolipoproteins from the circulation in, respectively the artery wall (Williams and Tabas, 1995; Tabas et al., 2007) and the BrM (Curcio et al., 2009; Al Gwairi et al., 1372 1373 2016) BrM. Indeed, proteoglycans may play an, as so far underestimated, role in 1374 regulating the complement response and the development of both AMD and 1375 atherosclerosis pathology (Tate et al., 1993; Toomey et al., 2018). Happonen and 1376 coworkers (2012) recently showed that small proteoglycans, such as PRELP, are 1377 regulators of the complement cascade (Happonen et al., 2012). Please note that 1378 choroidal cells produce PRELP (as suggested in the current study) and that this protein 1379 apparently accumulates in drusen. There are a few reports which have established the 1380 different patterns of distribution of large (Clark et al., 2011) and small proteoglycans (Keenan et al., 2012); the latter including biglycan, decorin, fibromodulin, lumican, 1381 1382 mimecan, opticin, and prolargin in post-mortem eye or vascular tissue. 1383 Both atherosclerosis and AMD patients may suffer from endothelial cell dysfunction. 1384 Accumulating evidence suggests that endothelial cell dysfunction may be the initiating 1385 step in atherosclerosis (Miteva et al., 2018). In their AMD studies, Schaumberg and 1386 coworkers provided epidemiological evidence that at least one marker for endothelial 1387 dysfunction and inflammation, sICAM-1 is linked to drusen formation and 1388 neovascularization (Schaumberg et al., 2007). Interestingly, higher levels of circulating 1389 endothelial cells (CECs), a biomarker for a diversity of systemic complications, including 1390 vascular disorders, were found in AMD patients compared to controls (Machalinska et 1391 al., 2011). 1392 While these studies focused on common risk factors and parallel development of drusen 1393 and plaques, the possibility that atherosclerosis plays a direct role in the development of 1394 AMD cannot be ruled out. Using FA, a slow filling of the choroidal capillaries over time 1395 has been observed in AMD patients (Pauleikhoff et al., 1990). This may be due to (a 1396 combination of) thickening of the BrM, a declining function of the RPE or by decreased 1397 atherosclerosis-driven perfusion of these capillaries. Reduced capillary blood flow could 1398 directly enhance the initiation of drusen or development of AMD. 1399 As with drusen, deposition of calcified mineral, that includes hydroxyapatite (Lee et al., 1400 2012) is associated with the formation of atherosclerotic plaques (Doherty et al., 2003). 1401 Such mineral is readily quantifiable using radiography and even serves as a marker for 1402 atherosclerosis. It has been reported that the presence of mineral in cardiovascular soft 1403 tissue can be used to predict mortality (Okuno et al., 2007; Kestenbaum et al., 2009), and 1404 morbidity of cardiovascular disease in various forms (Arad et al., 2000; Keelan et al., 1405 2001). The specific molecular mechanisms underlying mineral formation in such tissues 1406 remains to be fully elucidated. However, both AMD and atherosclerosis are associated 1407 with low grade inflammation in the respective affected tissues (Hansson et al., 2006; 1408 Kauppinen et al., 2016), and it has been proposed that soft tissue mineralization may be 1409 best conceptualized as a convergence of bone biology with inflammatory pathobiology 1410 (Doherty et al., 2003). 1411 8.3. Genetics and molecular biology. 1412 Early candidate gene association studies found an association between genetic variation 1413 in APOE in both AMD (Klaver et al., 1998; Toops et al., 2016) and atherosclerosis (Zhang 1414 et al., 2018), thereby implicating lipid metabolism and transport in both disorders. 1415 Genetic variations in apolipoproteins and complement factors showed strong 1416 associations with AMD and CVD conditions. For example, polymorphisms in the CFH and 1417 a number of other complement factor genes confer at-risk genotypes for AMD (Klein et 1418 al., 2005), whilst similar associations between complement C5 and the complement 1419 receptor 1 genes confer an increased risk of atherosclerosis (Hoke et al., 2012; de Vries 1420 et al., 2017). Of note, although the same genes may be frequently associated with both 1421 (or other) diseases, different alleles are frequently implicated in the associations found

1422 between these disorders. A well-known example is the APOE4 allele, that increases the 1423 risk of Alzheimer's disease, and perhaps atherosclerosis (Mahley, 2016), but is 1424 protective in age-related macular degeneration (Klaver et al., 1998). Indeed, these 1425 observations were confirmed and extended by large GWAS studies that implicated 1426 regulation of lipid metabolism, extracellular matrix remodeling and the immune system 1427 low-grade inflammation in both AMD and atherosclerosis (Fritsche et al., 2016; 1428 Schunkert et al., 2018). A recent study by the International AMD consortium explored 1429 the overlap between 34 AMD-associated loci with other complex diseases (Grassmann et 1430 al., 2017). Surprisingly, the authors found that an increased risk of AMD correlates with 1431 a reduced risk for cardiovascular disease. 1432 A key similarity between atherosclerotic plaque and drusen formation are the molecular 1433 components involved. Both types of deposit have a significant lipid component 1434 (including cholesterol and neutral fats) and mineral content, as described above. It has 1435 also been reported that drusen contains a number of proteins that are also common to 1436 atherosclerotic deposits (Mullins et al., 2000; Klein et al., 2005; Booij et al., 2010a). To 1437 gain information as to the degree of this overlap in proteins contributing to these 1438 pathologies we compared a dataset of 3196 proteins known to be present in 1439 atherosclerotic plaques from Bleijerveld and coworkers (Table 3) with our drusen data 1440 set, as shown in Figure 9a (Bleijerveld et al., 2013). The resultant Venn-diagram 1441 revealed that out the 89 drusen-associated proteins, 64 of these (72%) were also 1442 present in atherosclerotic plaques. Indeed, 50 out of 60 drusen proteins derived from 1443 blood are also present in atherosclerotic plaques (Figure 9b). Details of proteins found 1444 to be common to both plaques and drusen can be found in STable 11 and STable 11a. 1445 Closer inspection of proteins common to both atherosclerotic plaques and drusen as 1446 defined in this manuscript revealed a number of functional classes of protein in this 1447 group including apolipoproteins (APOA1, A2 and E), complement factors (C7, C8A, C8B, 1448 C8G and complement factor H), as well as lipid- and Ca²⁺-binding annexins (annexins-1, 1449 2, 5 and 6). Obviously, our analysis may not be fully comprehensive, since it is limited to 1450 the entries which are present in both database listings. Another limitation is that similar 1451 proteins still may originate from different sources. For example, the previously 1452 suggested presence of APOB as principal protein of LDL in both sub-RPE-BL deposits 1453 and cardiovascular plaques (Curcio et al., 2001) is missing from the current overlap,

since detailed investigation of the (presence and origin) of this lipoprotein (Li et al., 2005a) suggested that APOB isolated from BrM thickenings is (also) present in a distinct, non-LDL lipid profile. Consequently, it was suggested that APOB in BrM thickenings is made locally, while APOB in plaques is probably from systemic origin. Cytoskeletal proteins (actinin α 1, tubulin α 1c and tubulin β 3) as well as extracellular matrix proteins such as collagens (type 1 α 2, type 6 α 1, type 6 α 2 and type 8 α 1), tenascin C, microfibril-associated protein 4 and vimentin were found to be present in both BlamD deposits and atherosclerotic plaques (Fernandez-Godino et al., 2016; Pelisek et al., 2016). Analysis of proteins common to both plagues and drusen in biological processes revealed significant contribution of this group of proteins to other diseases and processes including various cancers, development of the vasculature, cell movement and AD (tauopathy and amyloidosis; see STable 11a). The involvement of these proteins in AD is particularly interesting as it is another disorder of which extracellular deposits are a feature (Figure 9c). Furthermore, drusen reside on the interface between the neural and cardiovascular system, so it may share properties of both types of atherosclerosis and Alzheimer's plaques (Booij et al., 2010a).

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9. Future directions and conclusions.

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1471 Our review of the literature and the qualitative and quantitative meta-analysis of retinal 1472 and blood transcriptomic and proteomic data all point in the same direction: proteins in 1473 drusen originate from multiple sources. Based on the data we have available, the largest 1474 number of protein contribution from a single source appears to be the blood. The 1475 second-most prominent source of number of specific proteins in drusen is from the RPE, 1476 while the contribution from the choroid and the photoreceptors appears to be relatively 1477 modest. However, the varying number of proteins cannot be directly translated to 1478 concentration. There is the possibility that a relatively small number of proteins 1479 contribute the bulk of proteins in drusen. 1480 How proteins get recruited to and retained in the sub-RPE-BL space is still not fully 1481 understood. *In vivo* and *in vitro* BrM conductance studies suggest that human proteins 1482 of average size, such as proteins of 53 kDa (source: NCBI), can readily diffuse through 1483 healthy BrM, while macromolecular migration through BrM is slower and/or limited 1484 (Curcio and Johnson, 2012). Moore and Clover found that proteins of 200 kDa could 1485 readily cross young BrM (Moore and Clover, 2001). More recent work suggested that the 1486 transport exclusion size limit in healthy young BrM can be as high as 180-500 kDa, well 1487 over the size of macro molecules like HDL (Hussain et al., 2010; Cankova et al., 2011). In 1488 our current study, we found that 9 out of 10 drusen proteins that are uniquely derived 1489 from the RPE had a Mw of less than 50 kDa (Table 6). Moreover, we took a random 1490 sample of 30 proteins from the RPE-IVS basal secretion dataset (Table 3), containing 1491 proteins which are likely to encounter BrM *in vivo*, and determined their average Mw: 1492 95 kDa). After taking three extremely large proteins out (APOB, AHNAK, and C4B; 1493 proteins that we did not identify in drusen in this study) that average dropped to 60 kDa 1494 (data not shown). Six of these 30 proteins are present in drusen (ALB, ANXA1, ANXA2, 1495 APOA4, APOE, ATP5F1B) and have a MW <66.5 kDa. Although the overall transport 1496 capability BrM decreases substantially in the AMD-affected and aging retina (Hussain et 1497 al., 2010; Cankova et al., 2011; Curcio and Johnson, 2012; Lee et al., 2015), older BrM 1498 was found to be still permeable to proteins in excess of 100 kDa (Moore and Clover, 1499 2001). Taken together, entrapment of proteins in the sub-RPE-BL space is unlikely to be 1500 due to size if single molecules. They might become entrapped by forming aggregates that 1501 are no longer capable of leaving through BrM, as was suggested for CFH (Nan et al.,

1502 2008; Nan et al., 2011; Nan et al., 2013). Therefore, drusen proteins, especially the ones 1503 that come from multiple sources, "meet, greet and stick" to form sub-RPE-BL space 1504 deposits. 1505 There are several ways proteins can interact in BrM to form larger aggregates: they can 1506 interact among themselves, with other lipids, proteins and/or mineral deposits, or stick 1507 to the ECM of BrM itself. These interactions may be enhanced by chemical modification, 1508 (Blaum et al., 2010) including oxidative damage and glycosylation of lipids, proteins and 1509 carbohydrates (Crabb et al., 2002; Hollyfield et al., 2010) and they may be further 1510 facilitated by the structure and dynamic nature of BrM (Booij et al., 2010a). Over time, 1511 several changes in BrM occur, that may hinder protein clearance from the sub-RPE-BL 1512 space. Remodeling of BrM ECM takes place, including proteoglycan changes and 1513 turnover, elastin changes and eventually mineralization takes place. BrM becomes laden 1514 with lipids to form a hydrophobic barrier ("lipid wall") and accumulates other debris 1515 (Curcio and Johnson, 2012). Consequently, the role of the structure and function of BrM 1516 and the chemical state of the sub-RPE-BL space may be even more important in sub-1517 RPE-BL space deposit formation than its exact protein composition. 1518 1519 An important source of entrapment of proteins in BrM may be the formation of HAP 1520 surfaces in the sub-RPE-BL space (Thompson et al., 2015). In our current study, at least 1521 30% of the 89 drusen proteins can bind to HAP. This percentage increases towards at 1522 least 50% if only the blood borne proteins are counted (data not shown). Thus, HAP 1523 readily binds a substantial number, but not all drusen proteins (Arya et al., 2018). 1524 The finding that blood proteins are seemingly the most important contributors to 1525 drusen formation provides a new target to prevent the initiation and propagation of sub-1526 RPE-BL space deposits. Reducing the concentration of blood proteins that interact with 1527 HAP may lead to a reduction of the source of drusen components and ultimately 1528 postpone, or potentially even stop, the progression to AMD. 1529 Finally, it is also important to mention that the non-specific interaction of proteins with 1530 HAP will also affect their ability to carry out their physiological function. For example, 1531 once CFH binds to the HAP surface it may not be able to regulate the alternative 1532 complement pathway. Therefore, this interaction with HAP could be a double whammy: 1533 it increases the bulk of sub-RPE-BL space deposits and stops the local protein function. It 1534 will be important to understand the role of the blood-derived proteins in the sub-RPE-

1535 BL space, if any. The study and potential modification of these interactions is now possible and could lead to intervention strategies through modified diet, 1536 1537 supplementation or through manipulation of retinal molecular or cellular processes. 1538 An important specific question in the context of this study that needs still to be resolved 1539 is how plasma proteins find their way into the sub-RPE-BL space. Apart from the 1540 mechanisms already described above (chemical modification of interacting 1541 biomolecules, dynamic structure and functional changes BrM, and HAP-binding) it is 1542 tempting to speculate that not only blood composition but also blood pressure plays a 1543 role. Why blood pressure? It was previously shown that a relationship exists between 1544 drusen location and choriocapillary pillars. Indeed, by investigating retina whole 1545 mounts, initially Friedman, and subsequently, Lengyel and coworkers concluded that 1546 drusen deposition is the result of a lower clearance at the choroidal intercapillary pillars 1547 (Friedman et al., 1963; Lengyel et al., 2004). Thus, in other words, higher clearance of 1548 sub-RPE-BL space debris corresponds with the vascular lumen, through which the blood 1549 flows and directly encounters BrM. Much in line with the reflections of Penfold and 1550 others (Penfold et al., 2001), we hypothesize that the pulsating blood pushes debris 1551 through endothelial fenestrations into BrM, through relatively open BrM pores; and at 1552 the same time, clears debris which was already present in BrM. One could compare that, 1553 by analogy, with the sea bringing and taking, wave after wave, debris to and from the 1554 beach. Changes in blood composition, choroidal endothelial cell compromise and rising 1555 blood pressure with age (Pinto, 2007) may negatively change the dynamics of this 1556 proposed "debris-exchange". 1557 Our review further underlines the importance of comparative studies between drusen 1558 deposition and atherosclerosis plaque formation. Clinical, (genetic) epidemiological 1559 pathobiological and molecular similarities between these two disorders have been 1560 highlighted previously (see section 8). Such similarities include that both are 1561 extracellular lipid/protein/mineral-based depositions that invoke a low-grade immune 1562 response leading to further disease. Several molecular similarities between drusen and 1563 plaques have also been described. We currently add the observation that most drusen 1564 (and plaque) components are blood-borne. Therefore, the genesis of drusen and plaques 1565 may be similar, and should be subject of further multidisciplinary studies.

1566 While studying the literature for this review, we have made a number of additional observations that may guide future research directions: First, while the number of 1567 1568 retinal (cell-type-specific) transcriptomics studies are large and proteomics information 1569 is emerging, there are very few proteomics studies on different types of (human) sub-1570 RPE-BL space deposits (types). For example, additional proteomics studies of hard 1571 versus soft drusen or macular versus peripheral drusen might improve our 1572 understanding of deposition formation in the sub-RPE-BL space and their association 1573 with different disorders or disease stages. Next, transcriptomics, proteomics, and 1574 immunohistochemical studies have their own conceptual and technical advantages and 1575 limitations. However, in the literature, the description of these strengths and 1576 weaknesses are not always clear and standardization is lacking. International 1577 agreements such as MIAME and MISFISHIE (minimum information specification for in situ hybridization and immunohistochemistry experiments) guidelines (Deutsch et al., 1578 1579 2008) are a step in the right direction, but must be seen as initial steps for further 1580 standardization. A few examples for illustration: How many confirmatory 1581 transcriptomics or proteomics studies should be performed before a definite subcellular 1582 assignment can be made? How do we define cellular specificity and cellular enrichment? How many drusen types should be screened and how many different antibodies should 1583 1584 be used before proteins are clearly assigned as drusen proteins (or as sub-types). When 1585 should we designate labeling drusen specific? Do we consider staining of the border of 1586 hydroxyapatite or drusen important; or is only the staining of the whole inner mass of 1587 drusen relevant? Given the heterogeneity of drusen: what is the exact location of the drusen under study and its appearance? Indeed, in line with recent similar calls by 1588 1589 Curcio and co-workers (Curcio, 2018a, b) we call here for better considerations, 1590 agreements and definitions of these issues. 1591 Last but not least, it will be interesting to understand whether drusen heterogeneity is a 1592 direct feature of a disease or a reflection of the change in the (micro-) environment that 1593 results in initiation and growth of the deposits. While drusen deposition clearly is a 1594 hallmark of AMD and is associated with a number of other diseases (Khan et al., 2016), 1595 its actual composition might reflect the disease state at the RPE/choroid interface more 1596 than (cause) the disease. The identification of why and not necessarily what proteins

- and lipids are deposited in the sub-RPE-BL space might therefore an important question
- 1598 to consider for future studies.

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Legends to Figures and (S)Tables.

Figure legends:

Figure 1. Heterogeneity of drusen. Imaging of drusen and drusen content with various clinical and laboratory methods. On color fundus images, the yellow spots identify drusen (A). On OCT image the elongated RPE is reflective. Drusen appear as homogeneous and hyper-reflective sub-RPE-BL space focal entities. (B); scale bar: 500 μm . On hematoxylin-eosin staining, drusen appears between the brown pigments of the RPE and the Bruch's membrane (C); scale bar: 10 μm . Note the inclusions without staining. Drusen contain numerous von Kossa positive spherule structures identifying hydroxyapatite spherules (D); scale bar: 10 μm . Auto-fluorescence of the more or less circular drusen is indicative of the protein and lipid accumulation. Auto-fluorescence of the RPE is more intense and yellowish. Auto-fluorescence of the BrM adjacent to the RPE is greenish (E); scale bar: 10 μm .

Figure 2 a-d. Functional molecular network analysis (4 networks). For all four networks: Molecular network analysis of physical or functional interactions between 89 drusen genes/proteins using Ingenuity. The knowledge database generated four most likely functional networks from the given input (drusen proteins). The functionalities in the figures are generated based on a combination of available molecular and cellular experimental data in human, mouse, rat, and in vitro data. In each network the circles, squares and other symbols represent proteins from a homologue from either human, rat or mouse. The systematic name of the gene/protein is printed on each symbol. Double circles in a single symbol denotes a group or /family of entries with a specific function and are sometimes introduced by the knowledge database to make networks possible or to simplify them. Solid lines represent strong physical or functional interaction between the entries, taken from published peer reviewed literature and/or transcriptomics and proteomics databases. Dotted lines represent weaker, associated relationships between the genes/proteins based on published peer reviewed experimental data (for example co-upregulation of expression in an in vitro experiment). The lines represent thus functionalities found in either, both, or all experiments on human, rat or mice (tissues) and in vitro findings. For example, the functional relationship between molecules A, B, C, could possibly be defined as follows A-functional mouse finding-B-functional human

finding-C. The underlying hypothesis is that the functionalities in human, mouse and rat are very similar. One can also generate networks of human (or rat or mice) functional data separately, but they are frequently quite similar, but less extensive.

In the first network (Figure 2a), clearly three functional clusters of closely related entries can be recognized: The complement cluster, the collagen cluster and the crystallin cluster. The second network (Figure 2b) is much more complicated and heterogeneous and is a network of genes and proteins related to development and genetic or ophthalmic disorders. The common theme of the third network (Figure 2c) is the immune response. Finally, the fourth and last network (Figure 2d) presents functional and structural relationships between entries involved in cell-cell interactions and systemic involvement.

Figure 3. Schematic overview of various strategies used for dataset curation. This figure shows several ways to curate (pre-existing) transcriptomics or proteomics datasets to form an improved, thorough or more specific dataset. For example, in (A) several datasets are merged into a cumulative new one by simply combining the datasets. The possible overlap is counted only once in the new merged database. One dataset was deleted, because it did not adhere to quality standard or had a different signature as the other ones. Strategy (B) has been published before (Booij et al., 2010b). In this case the original enriched RPE database contains 10% of the highest expressed genes in the cell. Some of the expressed genes in the RPE10% dataset overlap with the genes expressed in the adjacent tissues (photoreceptors and/or choroid). These "overlapping expressed genes" are therefore not specific of the RPE. Thus, to obtain a more specific (smaller) dataset, we discard of all the "overlapping expressed genes" in the RPE dataset, to obtain a highly enriched RPE dataset. Curation strategy C shows the breakdown of two datasets into desired subfamilies: The overlap between datasets X and Z is Y. Dataset Y can be used if overlap between X and Z is desired. Dataset X minus Y can be used to obtain unique entries from X (compared to set Z).

Figure 4. Scheme of the relationships between the respective transcriptomic and proteomic datasets used for quantitative studies. The curated drusen protein dataset represents 89 proteins known to be present in drusen/sub-RPE-BL space deposits (black box in the middle). These were compared with the entries present or produced

2461 uniquely from the neural side of drusen (photoreceptor/RPE neural source), and with 2462 entries uniquely from the systemic side of drusen (blood /choroid basal source). 2463 The "neural source" and "systemic source" merged data-sets each consist of non-curated 2464 datasets and curated datasets. The non-curated (pure) dataset contain, by virtue of their 2465 nature or previous curation in the literature) only entries from, respectively, the neural (RPE-ST; RPE-IVS) and systemic side (BL-SP1; BL-SP2) of drusen. The curated datasets 2466 2467 (cPRos; cPR-ET and cChor-ET)) contained, before curation, a number of entries 2468 expressed/present in both the neural and systemic side (see PRos; PR-ET and Chor-ET). 2469 Thus, we removed all overlapping expressed genes between the PRos; PR-ET and Chor-2470 ET datasets, to obtain unique datasets from both sides of drusen. 2471 2472 Figure 5. Potential contribution of neural and systemically expressed/present proteins 2473 to drusen formation. Venn diagram showing overlap between (A) neural RPE and 2474 photoreceptor-derived proteins, (B) systemically derived choroid and blood proteins 2475 and (C) drusen-associated proteins. 2476 2477 Figure 6. Hydroxyapatite spherules can retain proteins originating from blood in the sub-RPE-BL space. (A) Immunocytochemical labelling of histidine-rich glycoprotein 2478 2479 (HRG) using a specific anti-HRG primary antibody (green) on the surface of a HAP 2480 spherule labelled by LiCor680 (magenta); scale bar: 10 μm. (B) Binding of purified 2481 human HRG to HAP-coated magnetic beads. Binding assays were performed using 0.3 2482 mg beads per sample. HAP-beads were washed with 50 mM Tris, 140 mM NaCl, pH 7.4 2483 and incubated with 400 μ l of 0-1 μ M human HRG for an hour at room temperature. The 2484 protein-bound beads were washed with the same buffer twice followed by blocking with 2485 1% BSA for an hour. Rabbit anti-human HRG antibody (1:1000 dilution) and HRPconjugated anti-rabbit antibody (1:10000 dilution) were respectively used as primary 2486 2487 and secondary antibodies. Detection was done at 492 nm using o-phenylenediamine 2488 dihydrochloride (OPD, Sigma Aldrich) substrate. 2489 2490 Figure 7. A model for drusen formation. Top row (A-E) is adopted from the schematic 2491 diagram proposed for sub-RPE-BL space deposit formation by Thompson and colleagues 2492 (Thompson et al., 2015). (A) Healthy eyes show no sub-RPE-BL space deposit formation. 2493 (B) At Stage 1 lipid droplets are retained in the sub-RPE-BL space (black dot). (C) At

2494 Stage 2 mineralization occurs surrounding the lipid droplets (magenta ring). (D) At 2495 Stage 3 proteins bind to the HAP surfaces (blue ring). (E) At Stage 4 proteins and lipids 2496 start accumulating around the "seed" (yellow material). The bottom row (A'-E') shows 2497 morphological evidence for the prediction in the top row. (A') Retinal pigment 2498 epithelium forms a monolayer along the inner collagenous layer of the Bruch's 2499 membrane in healthy eyes (scanning electron microscopic image); scale bar: 10 μm. (B') 2500 Transmission electron micrograph of lipid droplets that accumulate in the sub-RPE-BL 2501 space; reproduced with permission from Curcio and Millican (Curcio and Millican, 2502 1999); scale bar: 2 μm. (C') Scanning electron microscopic identification of a single 2503 spherule located between the RPE basement membrane and the inner collagenous layer 2504 of Bruch's membrane; scale bar: 2 μm. (D') Immunofluorescent labelling of HRG (green) 2505 on the surface of a HAP spherule (magenta); scale bar: 2 µm. (E') An immunofluorescent 2506 labelling of complement factor H on a spherule surrounded by the autofluorescence of 2507 drusen (green) and RPE cells (yellow) (blue is DAPI staining the cell nuclei); scale bar: 2508 10 μm. 2509 2510 Figure 8. Schematic showing factors that are identified to contribute to mineralization of 2511 soft tissues and may contribute to HAP deposition in the sub-RPE-BL space. 2512 Abbreviations: ABCC6, ATP binding cassette subfamily C member 6; ANKH, ankylosis 2513 protein homolog; ATP, adenosine triphosphate; BrMP2, bone morphogenetic protein-2; 2514 BrMP2R, bone morphogenetic protein-2 receptor; BSP, bone sialoprotein; Ca, calcium; 2515 Cbfa-1, core-binding factor alpha-1; ENPP1, ectonucleotide 2516 pyrophosphatase/phosphodiesterase; Glu- and Gla-MGP, uncarboxylated- and 2517 carboxylated-matrix Gla protein; OPG, osteoprotegerin; OPN, osteopontin; Pi, inorganic 2518 phosphate; Pit-1, phosphate transporter-1; PPi, pyrophosphate; RANKL, receptor 2519 activator of nuclear factor kappa-B ligand; TNAP, tissue non-specific alkaline 2520 phosphatase. Figure adapted from (Ronchetti et al., 2013). 2521 Figure 9. Proteins present in atherosclerotic plaques and drusen. A. Venn diagram 2522 2523 showing 64 out of 89 drusen proteins overlap with the atherosclerotic plaque proteome, 2524 while 25 entries are unique to drusen in this comparison. B. Venn diagram showing 50 2525 out of 60 proteins (from the 89 drusen proteins) that come from blood (as unique 2526 source or shared with the PR/RPE) are actually present in atherosclerotic plaques. C.

2527 Venn diagram displaying the uniqueness and overlap of proteins between drusen (C.A), 2528 Alzheimer plaque proteins (C.B.) and atherosclerotic plaque proteins (C.C). The 2529 corresponding STable 11 and STable 11a present the corresponding entries in detail. 2530 2531 **Table Legends:** 2532 2533 Table 1. List of proteins present in the curated drusen dataset. We assembled a list of 89 2534 drusen proteins, mostly derived from the macular area, from the literature. For each 2535 entry the Gene symbol, Entrez gene name, location and type, human 2536 immunohistochemistry source and literature references are provided based on 2537 information found via the Ingenuity knowledge database (Qiagen, all rights reserved), relevant literature (PubMed searches) and other public databases, such as Genecard 2538 2539 (www.genecard.org) and DAVID (https://david.ncifcrf.gov/) 2540 Crabb 2002 (Crabb et al., 2002); Wang 2002 (Wang et al., 2010); Entries with *: although 2541 assigned to drusen by proteomics, IHC studies suggest a more likely protein location 2542 around or directly external from drusen. Further detailed investigation is warranted for these entries. **First detected in cynomolgus monkeys, afterwards in human drusen. 2543 2544 2545 Table 2. Summary of Ingenuity knowledge database core analysis of 89 proteins present 2546 in the curated drusen protein dataset. Summary of enriched motifs present in the 2547 dataset presented as top disease and biological functions, canonical pathways and 2548 discrete molecular networks. Note that these functional annotations types relate to 2549 either cellular (LRX/RXR/FXR activation; macrophages) or systemic (acute phase, 2550 atherosclerosis) entities. In the top disease and biological functions, we see that the 2551 dataset is enriched for hereditary disorders, ophthalmic disease, injury, metabolic diseases and developmental disorders. Finally, in the top functional or structural 2552 molecular networks, we find combinations of very basic functions (cancer and cellular) 2553 2554 to more specific pathobiological ones (ophthalmic and neurological disease etc. 2555 2556 Table 3. Summary of datasets used in this study and their respective functional clusters. 2557 Table displaying various datasets used in this study, along with their characteristics. In 2558 the first column, the result of the Ingenuity network analysis of drusen proteins is given 2559 in 4 significant molecular networks (N=1-4) corresponding to the networks shown in

Figure 2a-2d. Within these networks, six functional molecule clusters can be observed. For example, Network 1 (N1) contains 3 functional clusters: the complement (Network 1. cluster 1), the collagen (1.2) and the crystallines (heatshock) (1.3). Network 2 consist of 1 large cluster (2.4) being genetic and developmental ophthalmic disorders. Network 3 can be viewed as a cluster (# 5) of injury and inflammatory response and dermatological disease. Network 4 (N4) contains a cluster (4.6) of cell-to cell-signaling and systemic involvement. Column B gives the actual gene/protein names in these clusters. Column C states the overall functional annotation of these clusters. The first row of the Table from column D onward states the compartment of the datasets to be compared with drusen proteins in the functional clusters (within brackets, the number of entire in each dataset are given). In row 2 (acronym) from column D onward, the short and systematic acronym of each dataset is given. Row 3 (reference) contains from column D onward, the reference where the dataset can be found. Row 4 (methodology), from column D onward, contains the method by which the data were generated (transcriptomics, proteomics). Row five (source), from column D onward, contains the primary author who submitted the data or who can be contacted to obtain further information. The remaining boxes contains information which entries of the functional cluster are present both in drusen as well as in the transcriptomics or proteomics dataset(s). Combined analysis of the clusters in different datasets gives a qualitative idea from which cell type(s) drusen protein are derived.

Table 4. Summary of Ingenuity knowledge database core analysis of the curated photoreceptor gene expression (cPR-ET) dataset. Functional annotation of the curated and highly enriched photoreceptor cPR-ET database using the ingenuity knowledge database. The data driven top canonical pathways are highly relevant for photoreceptor function: Phototransduction pathway, glutamate receptor signaling, cholesterol biosynthesis and Wnt/Ca2+ signaling. The only surprise in our data-driven analysis could be the Huntinton disease signaling pathway. However, it has recently become clear that in Huntington's disease (HD), an inherited neurodegenerative disorder resulting in motor disturbances, cognitive and behavioral changes, deficits in retinal and visual processing function are significantly present (Coppen et al., 2018). Although we curated the PR database quite extensively, and thus selected for specific photoreceptor molecular signature and function, it is interesting to see that these motifs occur also in a

2593 number of other (top) diseases and functions, such as cancer, organismal injury, 2594 gastrointestinal disease, Hepatic disease and reproductive system disease. This may 2595 reflect the accumulating evidence that a substantial number of genetic or metabolic 2596 disease are also affect photoreceptor function. Similar to the canonical pathways and the 2597 biological motifs, the functional annotation of the photoreceptor selected molecular 2598 machinery apparently reflects a broad spectrum of biological and disease processes. 2599 2600 Table 5. Summary of Ingenuity knowledge database core analysis of the curated choroid, 2601 cChor-ET datasets. In this Table, we present the summary of the functional annotation of 2602 the choroid. Of course, the choroid is not a single tissue, but contains multiple cell types 2603 (endothelial cells, fibroblasts, macrophages, etc.) and the sample is inevitably 2604 contaminated with the blood. Within these limitations, data driven analysis of this 2605 specifically curated data set yielded a number of interesting enriched motifs, which 2606 indeed can be contributed to the choroid or blood: The canonical pathways indicate 2607 enriched immunological themes, such as the complement system, acute phase response 2608 signaling, and antigen-presenting cells, which is confirmed by several biological motifs 2609 (inflammatory disease and response, injury). Further, the canonical pathways generated, 2610 suggest an overlap between the molecular machinery of the choroid and atherosclerosis 2611 signaling. Indeed, in this manuscript, we devoted a whole section (8) to the 2612 pathobiological and molecular similarities 2613 between drusen and atherosclerotic plaques, and their –in time-associated diseases: 2614 AMD and atherosclerosis. Finally, a homology between hepatic function and choroid was 2615 observed. Indeed, there are a number of reports in the literature of cross-talk between 2616 liver and choroidal function, but that potential relationship remains to be elucidated. 2617 The final biological motifs are cancer and connective tissue disorders. Cancer, is of 2618 course very broad and frequently relates to blood vessel metabolism or (abnormal) cell 2619 division, while the connective tissue motif may relate to the action of local fibroblasts. 2620 The choroidal networks, show, again a very broad spectrum of molecular interactions, 2621 but this spectrum is quite distinct from the functional annotation of the photoreceptor 2622 networks presented in Table 5. 2623 2624 Table 6. Drusen proteins expressed or present in the PR/RPE and their characteristics. 2625 Overview and characteristics of ten drusen proteins, which most likely originate from

the neural side of drusen (namely PR and Chor). In the first column (A), general used abbreviations (according Gen bank) for gene/protein names are given. In column B, C, D respectively systematic Entrez number, cellular location and protein type corresponding to these proteins are presented. Column E and F contain the amino acid (aa) size and Molecular weight (Mw) of the proteins. Further the isoelectric point (pI; column G), the number of negative and positive charged aa residues (column H), the protein instability Index number (column I); the Alipathic index for solubility (I), and the GRAVY (hydrophobicity and hydrophilicity index). These are all standard characteristic of proteins which can be found in the Ingenuity database (Qiagen all right reserved) and public databases such as DAVID, (https://david.ncifcrf.gov), SWISS-prot (https://www.ebi.ac.uk/uniprot), Genecards (www.genecard.org) and/or the data shows that these entries apparently do not have specific characteristics, except perhaps for their ability to interact with one another, that could explain why they would get stuck in BrM as a drusen protein. We conclude that, if it is not the proteins that explain this, it must be the structure of BrM.

Supplementary Table Legends:

Table S1. List of 276 proteins present in the RPE-IVS dataset. For each entry the gene symbol, Entrez gene name, location and type are provided based on information found in the Ingenuity knowledge database.

Table S2. List of 170 proteins present in the RPE-ST dataset. For each entry the gene symbol, Entrez gene name, location and type are provided based on information found in the Ingenuity knowledge database.

Table S3. List of 412 proteins present in the Pros-EP dataset. For each entry the gene symbol, Entrez gene name, location and type are provided based on information found in the Ingenuity knowledge database.

Table S4. List of 995 proteins present in the BLP-SP1 dataset. For each entry the gene symbol, Entrez gene name, location and type are provided based on information found in the Ingenuity knowledge database.

2659 2660 Table S5. List of 262 HAP binding proteins in the BL-PHP blood proteome dataset. For 2661 each entry the gene symbol, Entrez gene name, location and type are provided based on 2662 information found in the Ingenuity knowledge database. 2663 2664 Table S6. List of 754 expressed genes present in the cPR-ET dataset. For each entry the 2665 gene symbol, Entrez gene IDs for human and mouse are provided. 2666 2667 Table S7. List of 848 expressed genes present in the cChor-ET dataset. For each entry the 2668 gene symbol, Entrez gene IDs for human and mouse are provided. 2669 Table S8. Annotation of 37 drusen proteins (out of 89) that may uniquely originate from 2670 2671 the blood. For each entry, the gene symbol. Entrez gene IDs for human and mouse are 2672 presented. 2673 2674 Table S8a Functional annotation of 37 drusen proteins that may originate from the 2675 blood. Combinations of genes/proteins in this group makes up specific functional categories associated with biological function or disease. 2676 2677 2678 Table S9. Annotation of 23 drusen proteins that may originate either from the neural or 2679 from the systemic side, using Ingenuity. For each entry its functional category, specific 2680 associated disease or function, p-value, gene names of associated proteins and number 2681 of proteins in each category are provided. 2682 2683 Table S9a Functional annotation of 23 drusen proteins that may originate from either 2684 the neural or the systemic side of drusen using Ingenuity. Combinations of 2685 genes/proteins in this group makes up specific functional categories associated with 2686 biological function or disease. 2687 2688 Table S10 Annotation of 19 drusen proteins of unclear origin. Entrez gene IDs for human 2689 and mouse are presented. 2690

2691	Table S10a Functional annotation of 19 drusen proteins of unclear origin using
2692	Ingenuity. For each entry its functional category, specific associated disease or function
2693	p-value, gene names of associated proteins and number of proteins in each category are
2694	provided.
2695	
2696	Table S11 List of 64 proteins common to both drusen and atherosclerotic plaques. For
2697	each entry, the gene symbol and Entrez Gene IDs for human and mouse and are
2698	provided.
2699	
2700	Table S11a Functional annotation of 64 proteins common to both drusen and
2701	atherosclerotic plaques. For each entry its functional category, specific associated
2702	disease or function, p-value, gene names of associated proteins and number of proteins
2703	in each category are provided.
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2723	Next page: Table 1: 89 drusen proteins used in this study.

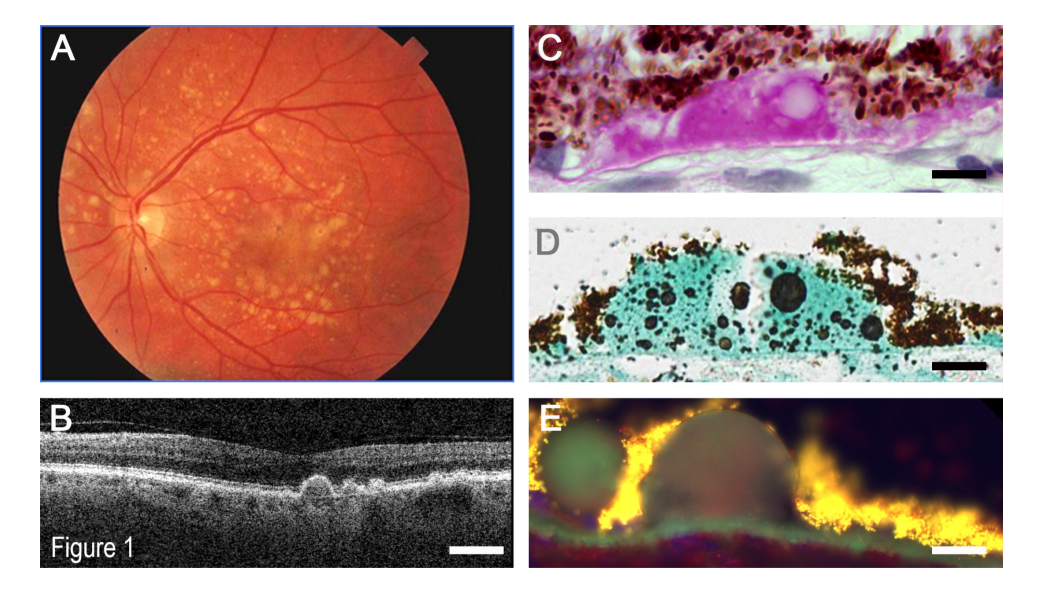
ID/Symbol	Entrez Gene Name	Location	Type(s)	Source	Human IHC ref
ACTB	Actin beta	Cytoplasm	other	(Crabb et al., 2002)	
ACTN1	Actinin alpha 1	Cytoplasm	transcription regulator	(Crabb et al., 2002)	
ALB	Albumin	Extracellular Space	transporter	(Crabb et al., 2002)	(Hollyfield et al., 2003)
ALDH1A1	Aldehyde dehydrogenase 1 family, A1	Cytoplasm	enzyme	(Crabb et al., 2002)	
AMBP	Alpha-1-microglobulin/bikufinbulin 5 drusennin precursor	Extracellular Space	transporter	(Crabb et al., 2002)	
ANXA1	Annexin A1	Plasma Membrane	enzyme	(Crabb et al., 2002)	(Rayborn et al., 2006)
ANXA2	Annexin A2*	Plasma Membrane	other	(Crabb et al., 2002)	(Crabb et al., 2002)*
ANXA5	Annexin A5	Plasma Membrane	transporter	(Crabb et al., 2002)	
ANXA6	Annexin A6	Plasma Membrane	ion channel	(Crabb et al., 2002)	(Crabb et al., 2002); (Rayborn et al., 2006)
APCS	Amyloid P component, serum	Extracellular Space	other	(Crabb et al., 2002)	(Mullins et al., 2000)
APOA1	Apolipoprotein A1	Extracellular Space	transporter	(Crabb et al., 2002)	(Mullins et al., 2000)
APOA4	Apolipoprotein A4	Extracellular Space	transporter	(Crabb et al., 2002)	
APOE	Apolipoprotein E	Extracellular Space	transporter	(Crabb et al., 2002)	(Mullins et al., 2000)
ATP5A1	ATP synth., H+ transp., mitochondr. F1 compl., alpha sub. 1, cardiac muscle	Cytoplasm	transporter	(Crabb et al., 2002)	
ATP5B	ATP synth., H+ transp., mitochondr. F1 compl., beta pp	Cytoplasm	transporter	(Wang et al., 2010)	
BFSP1	Beaded filament structural protein 1	Cytoplasm	enzyme	(Crabb et al., 2002)	
BFSP2	Beaded filament structural protein 2	Cytoplasm	other	(Crabb et al., 2002)	
BGN	Biglycan	Extracellular Space	other	(Crabb et al., 2002)	

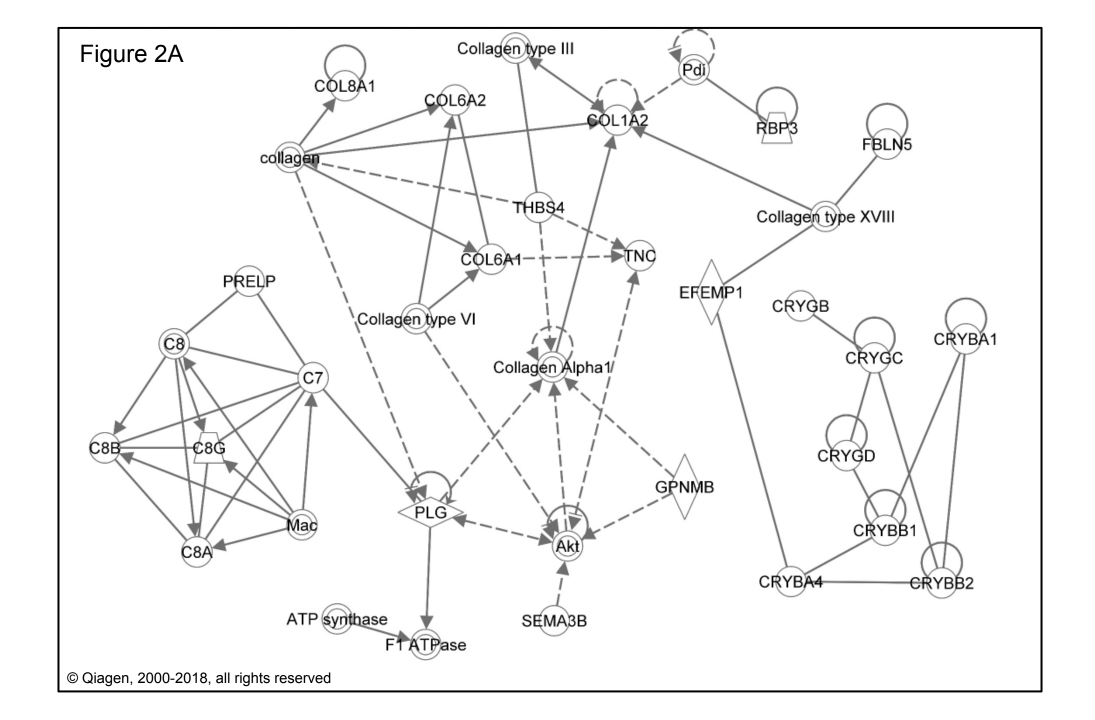
C7	Complement C7	Extracellular Space	other	(Crabb et al., 2002)	
C8A	Complement C8 alpha chain	Extracellular Space	other	(Crabb et al., 2002)	(Wang et al., 2010)
C8B	Complement C8 beta chain	Extracellular Space	other	(Crabb et al., 2002)	(Wang et al., 2010)
C8G	Complement C8 gamma chain	Extracellular Space	transporter	(Crabb et al., 2002)	(Wang et al., 2010)
CFH	Complement factor H	Extracellular Space	other	(Wang et al., 2010)	(Arya et al., 2018)
CKB	Creatine kinase B	Cytoplasm	kinase	(Crabb et al., 2002)	
CLU	Clusterin	Cytoplasm	other	(Crabb et al., 2002)	(Sakaguchi et al., 2002)
COL1A2	Collagen type I alpha 2 chain	Extracellular Space	other	(Crabb et al., 2002)	(Newsome et al., 1987)
COL6A1	Collagen type VI alpha 1 chain	Extracellular Space	other	(Crabb et al., 2002)	
COL6A2	Collagen type VI alpha 2 chain	Extracellular Space	other	(Crabb et al., 2002)	
COL8A1	Collagen type VIII alpha 1 chain	Extracellular Space	other	(Crabb et al., 2002)	
CRYAB	Crystallin alpha B*	Nucleus	other	(Crabb et al., 2002)	(De et al., 2007)*
CRYBA1	Crystallin beta A1	Other	other	(Crabb et al., 2002)	
CRYBA4	Crystallin beta A4	Other	other	(Crabb et al., 2002)	
CRYBB1	Crystallin beta B1	Other	other	(Crabb et al., 2002)	
CRYBB2	Crystallin beta B2	Other	other	(Crabb et al., 2002)	
CRYGB	Crystallin gamma B	Nucleus	other	(Crabb et al., 2002)	
CRYGC	Crystallin gamma C	Cytoplasm	other	(Crabb et al., 2002)	
CRYGD	Crystallin gamma D	Cytoplasm	other	(Crabb et al., 2002)	
CRYGS	Crystallin gamma S	Other	other	(Crabb et al., 2002)	
CTSD	Cathepsin D*	Cytoplasm	peptidase	(Crabb et al., 2002)	(Rakoczy et al., 1999)*

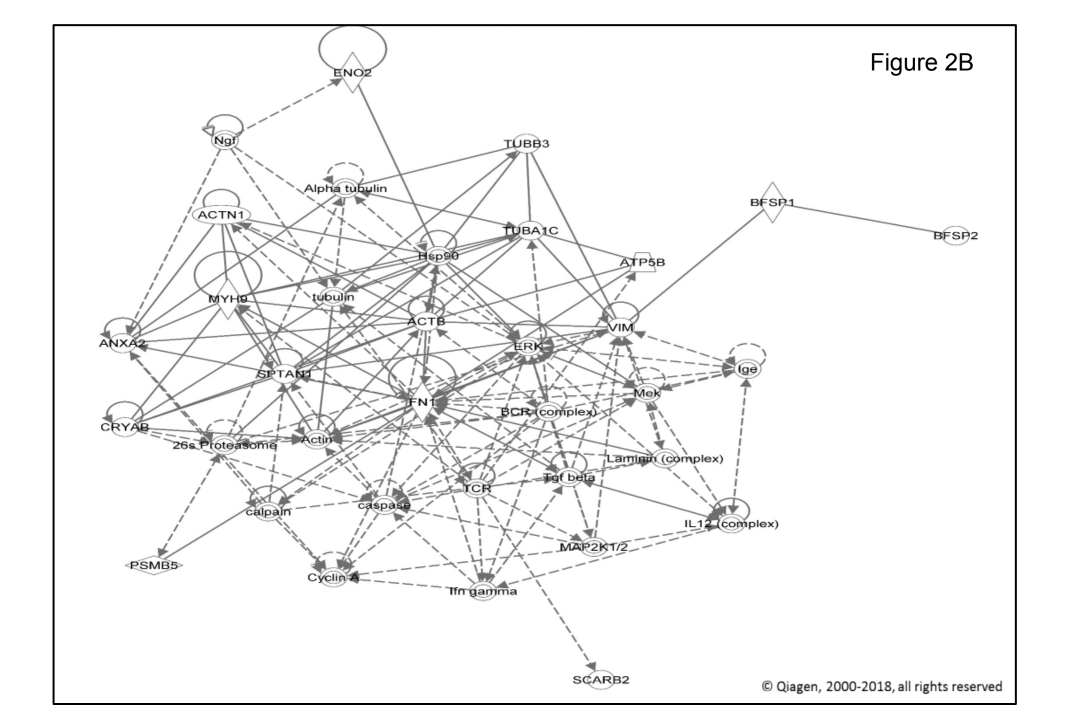
DIP2C	Disco interacting protein 2 homolog C	Other	other	(Crabb et al., 2002)	
EFEMP1	EGF containing fibulin like ECM protein 1	Extracellular Space	enzyme	(Crabb et al., 2002)	
ELN	Elastin*	Extracellular Space	other	(Crabb et al., 2002)*	
ENO2	Enolase 2	Cytoplasm	enzyme	(Wang et al., 2010)	
FBLN5	Fibulin 5	Extracellular Space	other	(Crabb et al., 2002)	(Mullins et al., 2000)
FGG	Fbrinogen gamma chain	Extracellular Space	other	(Crabb et al., 2002)	(Mullins et al., 2000)
FHAD1	Forkhead ass.phosphopept.bind. dom.	Other	other	(Wang et al., 2010)	
FN1	Fibronectin 1	Extracellular Space	enzyme	(Crabb et al., 2002)	(Newsome et al., 1987)
FRZB	Frizzled-related protein	Extracellular Space	other	(Crabb et al., 2002)	
GAPDH	Glyceraldehyde-3-phosphate dehydrogenase	Cytoplasm	enzyme	(Crabb et al., 2002)	
GPNMB	Glycoprotein nmb	Plasma Membrane	enzyme	(Crabb et al., 2002)	
HIST1H1E	Histone cluster 1 H1 family member e	Nucleus	other	(Crabb et al., 2002)	
HIST1H2BJ	Histone cluster 1 H2B family member j	Nucleus	other	(Crabb et al., 2002)	
HIST1H2B L	Histone cluster 1 H2B family member l	Nucleus	other	(Crabb et al., 2002)	
HIST2H2B E	Histone cluster 2 H2B family member e	Nucleus	other	(Crabb et al., 2002)	
HLA-DRA	Major histocompatibility complex, class II, DR alpha	Plasma Membrane	transmembrane receptor	(Wang et al., 2010)	
HRG	Histidine rich glycoprotein	Extracellular Space	other	(Kobayashi et al., 2014)**	Figure 6 and 7 (this study)
LAMB2	Laminin subunit beta 2	Extracellular Space	enzyme	(Crabb et al., 2002)	(Newsome et al., 1987)

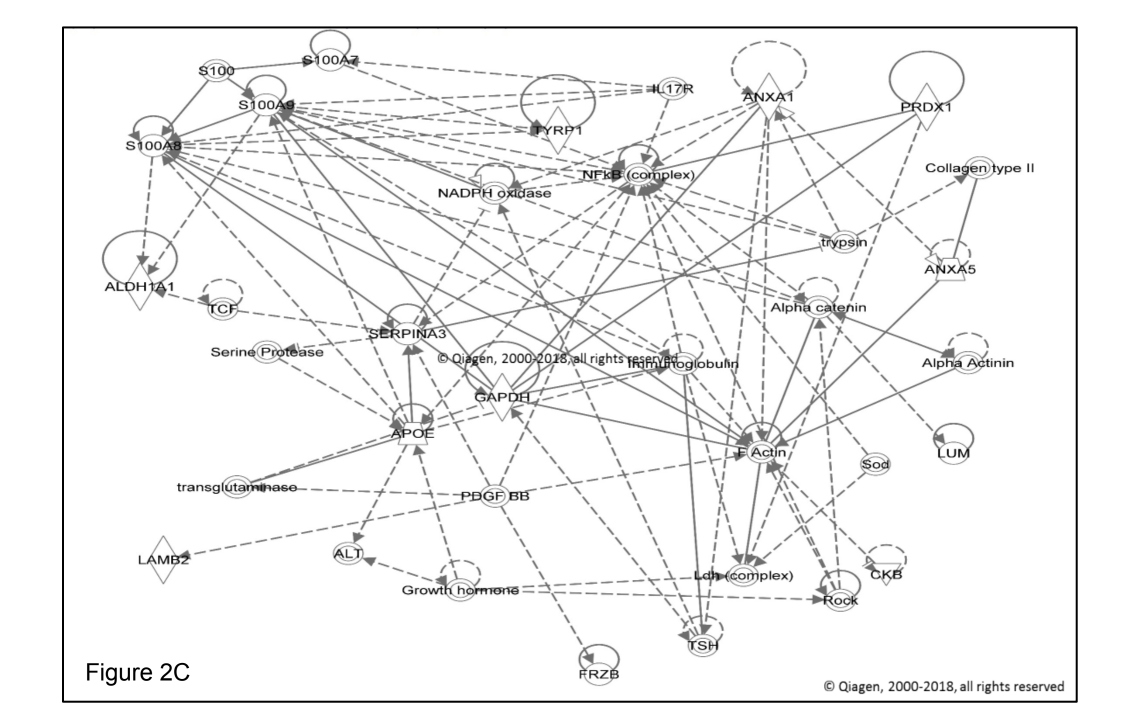
LTF	Lactotransferrin	Extracellular Space	peptidase	(Crabb et al., 2002)	
LUM	Lumican	Extracellular Space	other	(Crabb et al., 2002)	
MFAP4	Microfibril associated protein 4	Extracellular Space	other	(Crabb et al., 2002)	
MYH9	Myosin heavy chain 9	Cytoplasm	enzyme	(Crabb et al., 2002)	
OGN	Osteoglycin	Extracellular Space	growth factor	(Crabb et al., 2002)	
ORM1	Orosomucoid 1	Extracellular Space	other	(Crabb et al., 2002)	
PLG	Plasminogen	Extracellular Space	peptidase	(Crabb et al., 2002)	
PRDX1	peroxiredoxin 1	Cytoplasm	enzyme	(Crabb et al., 2002)	
PRELP	Pro, Arg rich end Leu rich repeat protein	Extracellular Space	other	(Crabb et al., 2002)	
PSMB5	Proteasome subunit beta 5	Cytoplasm	peptidase	(Crabb et al., 2002)	
RBP3	Retinol binding protein 3	Extracellular Space	transporter	(Crabb et al., 2002)	
RDH5	Retinol dehydrogenase 5	Cytoplasm	enzyme	(Wang et al., 2010)	
RGR	Retinal G protein coupled receptor	Plasma Membrane	G-protein coupled recept.	(Crabb et al., 2002)	
RNASE4	Ribonuclease A family member 4	Extracellular Space	enzyme	(Crabb et al., 2002)	
S100A7	S100 calcium binding proteA7	Cytoplasm	other	(Crabb et al., 2002)	(Crabb et al., 2002)
S100A8	S100 calcium binding protein A8	Cytoplasm	other	(Crabb et al., 2002)	(Crabb et al., 2002)
S100A9	S100 calcium binding protein A9	Cytoplasm	other	(Crabb et al., 2002)	(Crabb et al., 2002)
SAA1	Serum amyloid A1	Extracellular Space	transporter	(Crabb et al., 2002)	
SCARB2	Scavenger receptor class B member 2	Plasma Membrane	other	(Wang et al., 2010)	

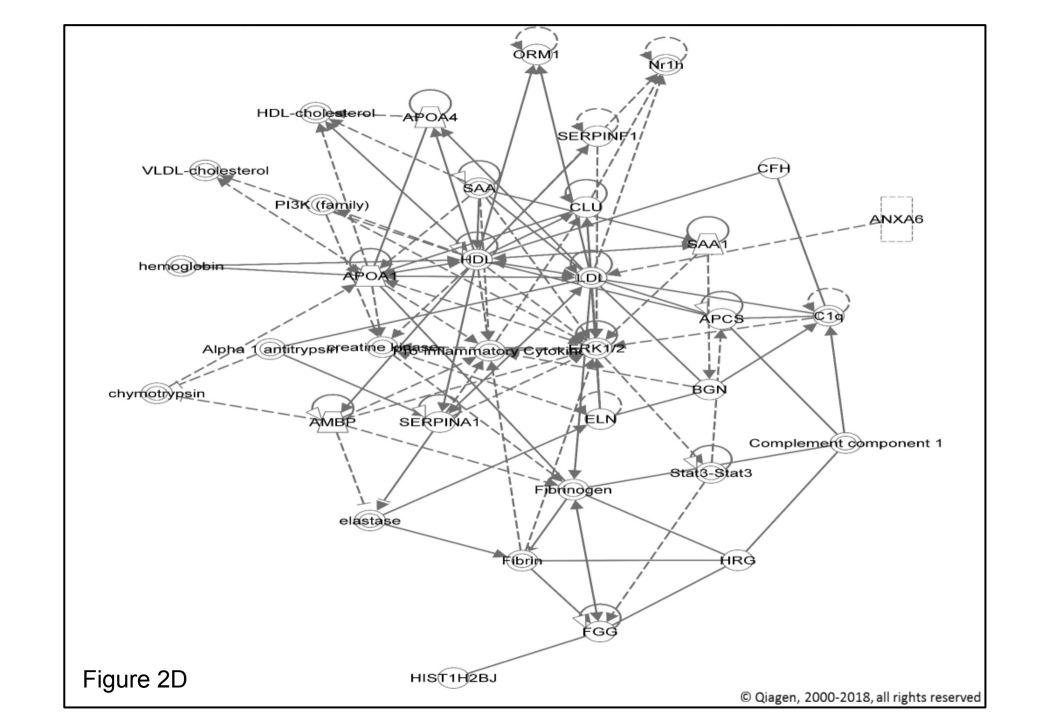
SEMA3B	Semaphorin 3B	Extracellular Space	other	(Crabb et al., 2002)	
SERPINA1	Serpin family A member 1	Extracellular Space	other	(Crabb et al., 2002)	
SERPINA3	Serpin family A member 3	Extracellular Space	other	(Crabb et al., 2002)	
SERPINF1	Serpin family F member 1	Extracellular Space	other	(Crabb et al., 2002)	
SPP2	Secreted phosphoprotein 2	Extracellular Space	other	(Crabb et al., 2002)	
SPTAN1	Spectrin alpha, non-erythrocytic 1	Plasma Membrane	other	(Crabb et al., 2002)	
THBS4	Thrombospondin 4	Extracellular Space	other	(Crabb et al., 2002)	
TIMP3	TIMP metallopeptidase inhibitor 3*	Extracellular Space	other	(Crabb et al., 2002)	(Kamei and Hollyfield, 1999)*
TNC	Tenascin C	Extracellular Space	other	(Crabb et al., 2002)	
TUBA1C	Tubulin alpha 1c	Cytoplasm	other	(Crabb et al., 2002)	
TUBB3	Tubulin beta 3 class III	Cytoplasm	other	(Crabb et al., 2002)	
TYRP1	Tyrosinase related protein 1	Cytoplasm	enzyme	(Crabb et al., 2002)	
VIM	Vimentin*	Cytoplasm	other	(Crabb et al., 2002)	(Johnson et al., 2003)*





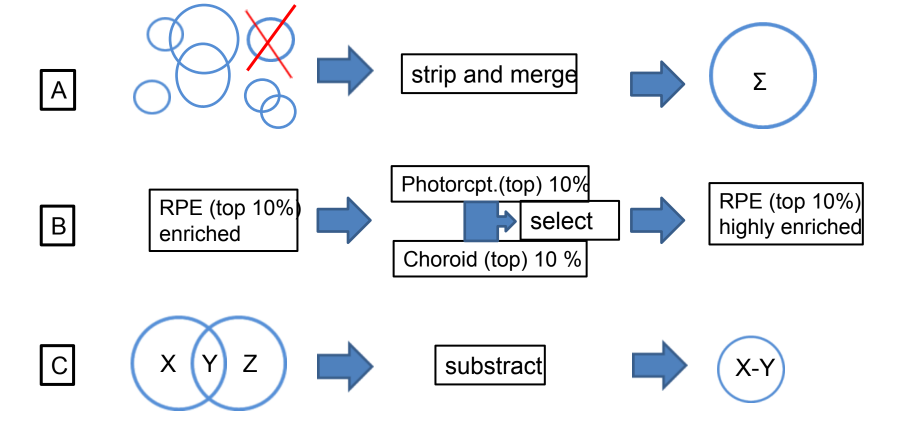


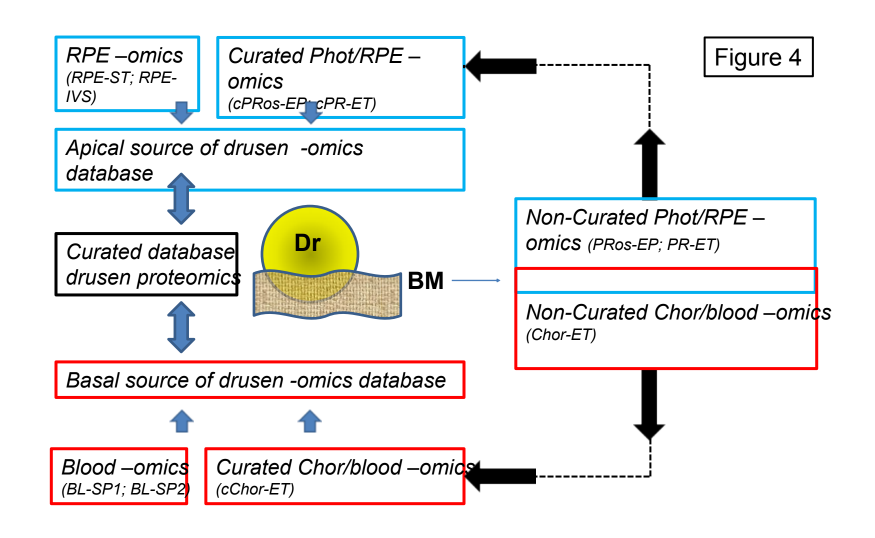


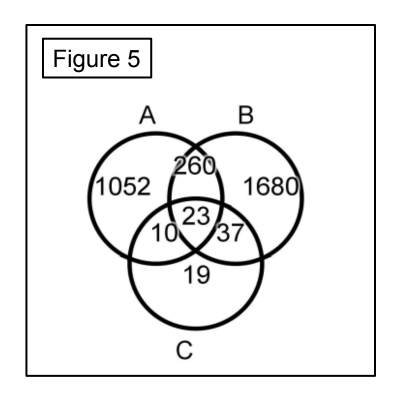


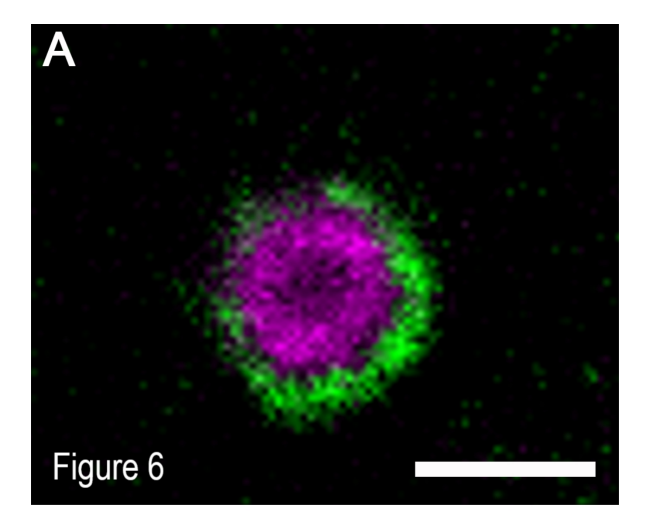
Database curation examples

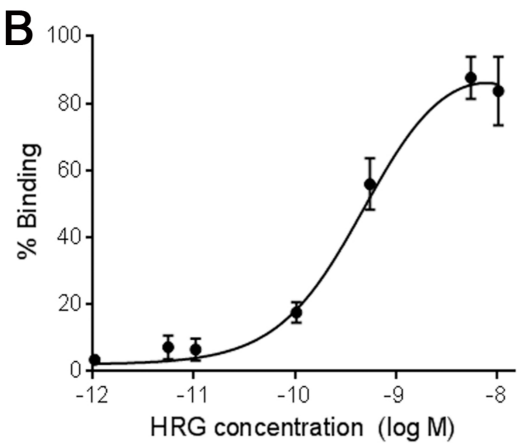
Figure 3

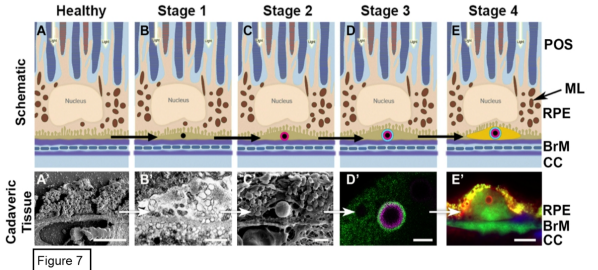


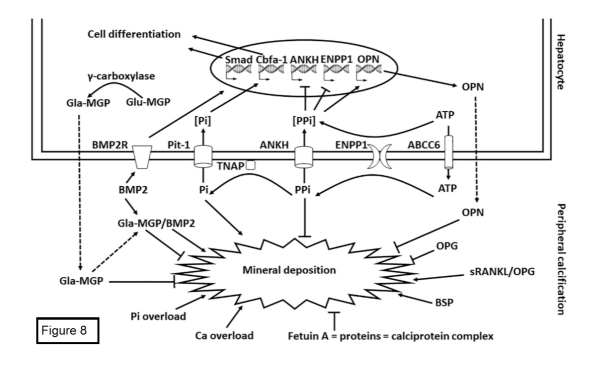


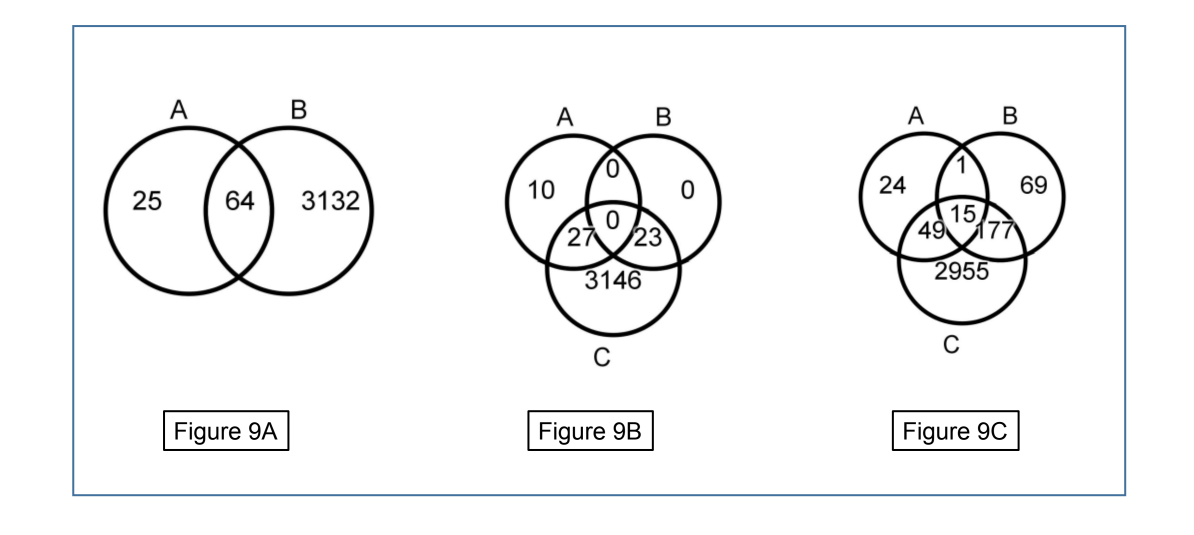












Ingenuity Pathway Analysis (IPA). Table 2.

Analysis Name: Table 2 Functional annotation 89 drusen proteins

Bergen et al.09:05 AM Analysis Creation Date: 2018-05-02

Build version: 470319M

Content version: 43605602 (Release Date: 2018-03-28)

Top Canonical Pathways

Name	p-value	Overlap
Acute Phase Response Signaling	7,51E-15	8,2 % 14/170
LXR/RXR Activation	2,29E-12	9,1 % 11/121
FXR/RXR Activation	9,39E-11	7,9 % 10/126
Atherosclerosis Signaling	2,35E-09	7,1 % 9/127
IL-12 Signaling and Production in Macrophages	1,42E-07	5,5 % 8/146

Top Diseases and Bio Functions

Diseases and Disorders

Name	p-value	#Molecules
Hereditary Disorder	1,04E-04 - 3,14E-19	51
Ophthalmic Disease	1,04E-04 - 3,14E-19	37
Organismal Injury and Abnormalities	1,19E-04 - 3,14E-19	88
Metabolic Disease	9,39E-05 - 4,75E-14	47
Developmental Disorder	1,04E-04 - 9,87E-14	34

Molecular and Cellular Functions

Name	p-value	#Molecules
Cellular Movement	1,18E-04 - 2,86E-16	44

Summary of Analysis - Table 2 Final annotation 89 drusen proteins 2018 - 2018-05-02 09:05 AM

Cell-To-Cell Signaling and Interaction	1,18E-04 - 1,74E-12	43
Lipid Metabolism	9,71E-05 - 1,02E-10	27
Molecular Transport	1,09E-04 - 1,02E-10	33
Small Molecule Biochemistry	9,71E-05 - 1,02E-10	27
Physiological System Development and Function	n value	#Molecules
Physiological System Development and Function	p-value	#Molecules
Name	p-value 1,04E-04 - 3,33E-17	#Molecules 30
	•	
Name	•	

1,04E-04 - 3,33E-17

1,08E-04 - 3,33E-17

Top Networks

Organismal Development

Tissue Development

ID Associated Network Functions	Score
1.Cancer, Connective Tissue Disorders, Organismal Injury and Abnormalities	44
2.Developmental Disorder, Ophthalmic Disease, Organismal Injury and Abnormalities	30
3.Cell-To-Cell Signaling and Interaction, Hematological System Development and Function, Lipid Metabolism	27
4.Cell-To-Cell Signaling and Interaction, Cellular Assembly and Organization, Neurological Disease	25
5. Neurological Disease, Infectious Diseases, Respiratory Disease	22

53

52

Ingenuity Pathway Analysis (IPA). Table 4.

Analysis Name: Table 4 Summary photoreceptor core annotation analysis (745); Bergen et al.

2018-05-07 10:07 PM Analysis Creation Date: 2018-05-07

Build version: 470319M

Content version: 43605602 (Release Date: 2018-03-28)

Top Canonical Pathways

Name	p-value	Overlap
Phototransduction Pathway	7,98E-11	28,3 % 15/53
Huntington's Disease Signaling	2,04E-04	8,0 % 20/250
Glutamate Receptor Signaling	4,79E-04	14,0 % 8/57
Superpathway of Cholesterol Biosynthesis	1,87E-03	17,9 % 5/28
Wnt/Ca+ pathway	4,20E-03	11,1 % 7/63

Top Diseases and Bio Functions

Diseases and Disorders

Name	p-value	#Molecules
Cancer	1,31E-02 - 6,87E-28	680
Organismal Injury and Abnormalities	1,33E-02 - 6,87E-28	685
Gastrointestinal Disease	1,21E-02 - 4,26E-22	629
Hepatic System Disease	2,47E-03 - 8,92E-15	474
Reproductive System Disease	1,27E-02 - 3,21E-08	420

Molecular and Cellular Functions

Name	p-value	#Molecules
Cellular Assembly and Organization	1,28E-02 - 1,06E-08	165

Summary of Analysis - photoreceptor core annotation analysis; Bergen et al. (745) - 2018-05-07

Cellular Function and Maintenance	1,31E-02 - 1,06E-08	185
Cell Death and Survival	1,32E-02 - 3,71E-06	240
Cell Morphology	1,31E-02 - 5,38E-06	149
Cell-To-Cell Signaling and Interaction	1,16E-02 - 8,21E-06	61

Physiological System Development and Function	p-value	#Molecules
Name	1,25E-02 - 3,12E-06	88
Organ Development		
Tissue Development	1,28E-02 - 3,12E-06	129
Visual System Development and Function	1,16E-02 - 3,12E-06	30
Nervous System Development and Function	1,28E-02 - 8,07E-06	161
Tissue Morphology	1,28E-02 - 9,43E-06	92

Top Networks

ID Associated Network Functions	Score
1.Cellular Assembly and Organization, Cellular Function and Maintenance, Molecular Transport	50
2.Molecular Transport, RNA Trafficking, Behavior	47
3.Developmental Disorder, Neurological Disease, Cellular Assembly and Organization	47
4.Molecular Transport, RNA Trafficking, Connective Tissue Development and Function	42
5.Developmental Disorder, Hereditary Disorder, Organismal Injury and Abnormalities	42

Ingenuity Pathway Analysis (IPA). Table 5.

Analysis Name: Table 5 Summary Functional Annotation Choroid- Bergen et al

Analysis Creation Date: 2018-05-07

Build version: 470319M

Content version: 43605602 (Release Date: 2018-03-28)

Top Canonical Pathways

Name	p-value	Overlap
Antigen Presentation Pathway	7,58E-12	36,8 % 14/38
Atherosclerosis Signaling	3,26E-11	18,0 % 23/128
Hepatic Fibrosis / Hepatic Stellate Cell Activation	3,88E-11	14,7 % 28/191
Acute Phase Response Signaling	5,19E-10	14,5 % 25/172
Complement System	1,90E-09	31,6 % 12/38

Top Diseases and Bio Functions

Diseases and Disorders

Name	p-value	#Molecules
Cancer	1,44E-06 - 4,74E-32	730
Organismal Injury and Abnormalities	1,48E-06 - 4,74E-32	746
Inflammatory Response	1,09E-06 - 1,87E-21	263
Connective Tissue Disorders	1,48E-06 - 1,46E-17	192
Inflammatory Disease	4,57E-08 - 1,46E-17	174

Molecular and Cellular Functions

Name	p-value	#Molecules
Cellular Movement	1,26E-06 - 1,81E-32	257

Summary of Analysis -

Cell Death and Survival	1,47E-06 - 4,17E-21	323
Cell-To-Cell Signaling and Interaction	1,50E-06 - 9,12E-15	218
Cellular Development	1,49E-06 - 1,14E-14	326
Cellular Function and Maintenance	8,75E-07 - 3,23E-12	274

Physiological System Development and Function	p-value	#Molecules
Name	9,30E-07 - 1,46E-28	192
Cardiovascular System Development and Function		
Organismal Development	1,47E-06 - 1,38E-24	325
Immune Cell Trafficking	1,29E-06 - 1,10E-23	158
Hematological System Development and Function	1,29E-06 - 3,90E-23	236
Organismal Survival	5,54E-08 - 1,79E-22	244

Top Networks

ID Associated Network Functions	Score
1.Organ Morphology, Organismal Injury and Abnormalities, Renal Atrophy	49
2.Organismal Injury and Abnormalities, Skeletal and Muscular Disorders, Developmental Disorder	41
3.Molecular Transport, Nucleic Acid Metabolism, Small Molecule Biochemistry	38
4. Tissue Development, Cellular Movement, Hair and Skin Development and Function	37
5.Cell Cycle, Gene Expression, Cellular Growth and Proliferation	36



Authors' statement

All authors have seen and approved the final version of the manuscript being submitted. They warrant that the article is the authors' original work, hasn't received prior publication and isn't under consideration for publication elsewhere.

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