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3	Evolution of Human-specific Alleles Protecting Cognitive Function of Grandmothers							
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Summary (250 Words)

27 Late-onset Alzheimer's Disease (LOAD) pathology is rare in our closest living evolutionary 28 relatives (chimpanzees), which also express much lower microglial levels of CD33(Siglec-3)-a 29 myelomonocytic receptor inhibiting innate immune reactivity by extracellular V-set domain 30 recognition of sialic acid(Sia)-containing "self-associated molecular patterns" (SAMPs). We 31 earlier showed that V-set domain-deficient CD33-variant allele, protective against LOAD, is 32 derived and specific to hominin-lineage. We now report that CD33 also harbors multiple hominin-33 specific V-set domain mutations and explore selection forces that may have favored such genomic 34 changes. N-glycolylneuraminic acid (Neu5Gc), the preferred Sia-ligand of ancestral CD33 is 35 absent in humans, due to hominin-specific, fixed loss-of-function mutation in CMAH, which 36 generates CMP-Neu5Gc from its precursor, CMP-N-acetylneuraminic acid (Neu5Ac). Extensive 37 mutational analysis and MD-simulations indicate that fixed change in amino acid 21 of hominin 38 V-set domain and conformational changes related to His45 corrected for Neu5Gc-loss by switching to Neu5Ac-recognition. Considering immune-evasive "molecular mimicry" of SAMPs 39 40 by pathogens, we found that human-specific pathogens Neisseria gonorrhoeae and Group B 41 Streptococcus (affecting fertility and fetuses/neonates respectively) selectively bind huCD33 and 42 this binding is significantly impacted by amino acid 21 modification. Alongside LOAD-protective 43 CD33 alleles, humans harbor additional, derived, population-universal, cognition-protective 44 variants absent in "great ape" genomes. Interestingly, 11 of 13 SNPs in these human genes 45 (including CD33), that protect the cognitive health of elderly populations, are not shared by 46 genomes of archaic hominins: Neanderthals and Denisovans. Finally, we present a plausible 47 evolutionary scenario to compile, correlate and comprehend existing knowledge about huCD33 48 evolution and suggest that grandmothering emerged in humans.

- 49
- Keywords (up to 10 words): Siglec3/CD33, pathogens, sialic acids, archaic genome, molecular
 dynamics simulation, phylogenetic analysis, menopause, grandmother.
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Introduction

54 In keeping with the fundamental importance of reproduction for the process of biological evolution 55 via natural selection, loss of fecundity generally coincides with the end of lifespan in almost all 56 species studied to date. Humans and certain toothed whales like orcas are so far the only mammals 57 known to manifest prolonged post-reproductive lifespans under naturalistic conditions [1–6]. One 58 current explanation for such prolonged post-reproductive survival is late-life kin selection of 59 grandmothers and other elderly caregivers of helpless young; apparently contrary to the concept 60 of "antagonistic pleiotropy", which posits that natural selection does not operate in late-life [7, 8]. 61 An interesting exception is a human-specific derived allele of CD33 associated with direct or 62 indirect protection against late-onset Alzheimer's Disease (LOAD) [9]. Furthermore, we noted 63 that humans harbor additional examples of such derived, population-universal gene variants that 64 directly or indirectly impact late-life cognitive decline, which were not found in other "great ape" 65 genomes. This was considered as genomic evidence for the evolution of human postmenopausal longevity [10]. Here we further explore the human-specific, derived alleles of genes that protect 66 67 against late-life cognitive decline, and ask when and how these emerged in hominins?

68 In vertebrates, glycan-binding proteins of the immunoglobulin (Ig) superfamily called sialic acid (Sia)-binding Ig-like lectins (Siglecs) form a major component of the immune system 69 70 [11]. As the name indicates, Siglecs recognize Sias on cell surface or secreted glycoproteins and 71 glycolipids. Siglec-3, commonly known as CD33, is the eponymous member of the rapidly 72 evolving subgroup of Siglecs called CD33-related Siglecs (or CD33rSiglecs) [12, 13]. In contrast, 73 other Siglecs (Siglecs 1, 2, 4 and 15) show evolutionary conservation [14]. CD33 is a type-I 74 transmembrane protein with an amino terminal Ig-like V-set domain followed by one Ig-like C2-75 set domain proximal to the transmembrane region [13]. Its cytoplasmic tail contains 76 immunoregulatory signaling motifs called immunoreceptor tyrosine-based inhibitory motif 77 (ITIM)s, which upon ligand binding to the extracellular V-set domain, undergo phosphorylation 78 and recruit effector molecules like tyrosine phosphatases, SHP-1/2, which inhibit the cellular 79 immune response. Human CD33 (huCD33) binds α 2-3- and α 2-6-linked N-acetylneuraminic acid 80 (Neu5Ac), the predominant Sia in humans, associated either with N- and O-glycosylated 81 molecules or sialylated glycolipids (gangliosides). HuCD33 undergoes alternative splicing, 82 resulting in two isoforms - full length CD33M containing the ligand binding V-set domain and 83 truncated D2-CD33 (or CD33m) lacking this domain [15]. The elimination of the terminal V-set

domain is mediated through differential splicing affected by two co-inherited single nucleotide polymorphisms (SNPs) at positions rs3865444 in huCD33 promoter and rs12459419 located within exon 2 [16]. The two isoforms, CD33M and D2-CD33, differ not only in their molecular weights, but also in their cellular localization and functionality which are associated with Siainteracting V-set domain [9, 12, 17].

89 HuCD33 is extensively studied for its role in different immune responses, under both 90 normal and pathophysiological conditions including cancers [16, 18, 18–21]. Furthermore, the 91 microglial expression of CD33 is linked with neurological pathologies like LOAD. Incidence of 92 LOAD has been strongly associated with varied expression of CD33 isoforms in the brain of 93 affected individuals [20, 21], where the LOAD-protective CD33 allele increases the ratio of D2-94 CD33 isoform relative to CD33M. CD33 is reported in almost all vertebrates, including nonhuman 95 primates [6, 14]. While there is often high similarity in the sequence and overall genomic location, 96 CD33 has undergone various species-specific changes. For example, murine CD33 which shows 97 about 54% identity with huCD33 V-set and 72% identity with C2 domain, has markedly different 98 Sia-binding and cellular expression patterns from human CD33 protein [22]. CD33 expression has 99 greatly diverged in humans even in comparison to our closest living evolutionary relatives, the 100 great apes. Examination of CD33 in peripheral blood showed significantly increased production 101 of CD33M in human monocytes relative to those of chimpanzees [9]. Furthermore, the abundance 102 of CD33 was also markedly higher in the human brain. Interestingly, although LOAD-associated 103 neurological pathologies, for example, buildup of A^β proteins, hyperphosphorylated tau proteins 104 as neurofibrillary tangles, have been observed in aged nonhuman primate brains, AD has largely 105 been regarded as a uniquely human disease [23, 24]. Interspecies variations in CD33 have also 106 been studied in other apes like gorilla and bonobo, in comparison to huCD33 [25].

107 The presence of two physiologically significant isoforms, their distinct cellular localization and 108 association with uniquely human pathologies like LOAD have made huCD33 a target of much 109 evolutionary interest. The Sia-binding V-set domain of CD33rSiglecs including CD33 itself show 110 high sequence variability amongst different species, often making it difficult to identify their 111 orthologs. The selective pressure for this accelerated evolution of the V-set domains has been 112 attributed to evasion of infectious pathogens that exploit these human innate receptors. The 113 surfaces of each vertebrate cell are layered with tens to hundreds of million Sia-terminating 114 glycans, forming as "self-associated molecular patterns" (SAMPs), which prevent erroneous

115 activation of innate immune responses against the body's own cells [26]. However, several human 116 pathogens e.g., Neisseria gonorrhoeae, Neisseria meningitidis, Haemophilus influenzae, E. coli 117 K1, Group B Streptococcus, and Trypanosoma cruzi cloak themselves with sialoglycans, 118 effectively mimicking host SAMPs, and thereby avoiding the immune response [27]. Conversely, 119 other infectious agents like influenza virus recognize SAMPs and utilize them as receptors to 120 initiate binding and subsequent infections [28]. CD33 has also been shown to interact with 121 Hepatitis B viral surface sialoglycans, thereby impacting its pathogenesis [29]. SAMPs and their 122 interacting partners, Siglecs (primarily the V-set domains) are therefore continually evolving to 123 maintain their distinctive "self-recognition" properties, while also avoiding exploitation by Sia-124 cloaked pathogens and parasites – a powerful example of the "Red Oueen Effect" [30].

125 In this work, with a focus on CD33, a post-reproductive cognitive health associated human protein, 126 we attempt to explore the evolutionary pressures that selected for unique changes in huCD33. 127 Using human-specific pathogens like Neisseria gonorrhoeae, Group B Streptococcus and E. coli 128 K1, we demonstrate differential impact of these mutations on the bacterial interactions with 129 huCD33. We also determine the effect of these mutations on huCD33-sialoglycan binding and 130 identify that the amino acid at position 21 within the V-set domain plays a critical role in Sia-131 specificity of human and chimpanzee CD33. Furthermore, we extend our study to archaic hominin 132 genomes and show that the human-specific CD33 mutations (except the presence of truncated 133 isoform) are shared evolutionary changes of human, Neanderthal and Denisovan common 134 ancestor. We also expanded our analysis to include other human-specific derived genomic changes 135 associated with cognitive health of post-reproductive human grandmothers and other elderly 136 caregivers. Finally, we draw an evolutionary scenario to connect the current knowledge of CD33 137 sialoglycan recognition and pathogen engagement to propose a role for the infectious pathogens 138 as key selective agents in human-specific CD33 evolution, generating new alleles protective 139 against infections, that could secondarily have come under selection for their protective effects 140 against cognitive pathologies like LOAD.

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Results

143 Sequences of human CD33 extracellular domains show many changes distinct from closely 144 related great apes. Previous investigations have identified unique properties of huCD33 that 145 influence the functionality of this molecule in humans. The presence of a huCD33 V-set truncated

146 isoform as well as its overall expression difference in microglia has been associated with the 147 protection against the occurrence of neurological pathologies like LOAD in humans. Like other 148 CD33rSiglecs, CD33 immunomodulatory roles depend both on its ligand-interacting extracellular 149 domains and signaling motif-containing cytoplasmic tail. To gain a comprehensive understanding 150 of different CD33 domain variations, we compared the amino acid residues of full-length CD33 151 from human and related nonhuman primates including chimpanzee, gorilla and bonobo (Figure 152 1A). While the regions encoding the C2-set domain and cytoplasmic tail are highly conserved, the 153 amino acid residues within huCD33 V-set domain differ from their nonhuman counterparts in as 154 many as 10 positions. Since different amino acid residues in Sia-binding V-set domain could 155 potentially impact huCD33-sialoglycan interactions and subsequent downstream signaling 156 pathways, we further examined the overall frequency of these changes (Figure 1B). We analyzed 157 human sequences from the 1000 Genome database [31] and compared them with 44 gorilla, 59 158 chimpanzee and 10 bonobo sequences [32-34]. Most of these amino acid residues (except at 159 positions 66 and 148) are conserved in all the great apes and appeared to have changed only in the 160 human lineage. Interestingly, the amino acid residues at positions 66 and 148 in huCD33 are 161 isoleucine and leucine respectively, similar to CD33 of chimpanzee and bonobo. The 162 corresponding amino acids in its more distant evolutionary relative, gorilla, are phenylalanine 163 (Phe) (at position 66) and valine (at position 148). The presence of the same amino acid in human, 164 chimpanzee and bonobo at these positions suggests a more ancient occurrence of these two 165 changes, possibly prior to the divergence of chimpanzee about 6-8 million years ago (mya). 166 Previously it has been shown that the two linked SNPs, resulting in the splicing of the V-set 167 truncated isoform represent a derived evolutionary modification of the CD33 proteins in humans 168 and are absent in chimpanzees [9].

To further understand the selection pressure, we calculated the nonsynonymous to synonymous substitution rate ratio (omega, $\omega = d(N)/d(S)$) for the CD33 V-set domains of human and other great apes. The omega value of CD33 V-set domain is greater than >1 (ω =1.49) which reflects Vset domain evolution under positive selection. Subsequently, we also analyzed the Ka/Ks ratios of exon 2 sequences in every species. Except for gorilla, the other two great apes (chimpanzee and bonobo) showed Ka/Ks ratios greater than one indicating that high Ka/Ks ratio of exon 2 is not an accidental event but an evolutionary phenomenon. Taken together, these results demonstrate that 176 CD33 in humans has been rapidly evolving possibly under positive selection, distinct from its177 orthologs in the great apes.

178 Archaic Neanderthal and Denisovan genomes share most human CD33 protein changes, 179 except for the SNPs for the LOAD-protective allele. Divergence of humans from other ancient 180 hominin lineage such as Neanderthals and Denisovans has been estimated to date back 181 approximately 0.5 mya [35]. Although full length CD33 itself is an ancient molecule, we noted 182 that the AD-protective CD33 truncated isoform is recently derived in humans, postdating our 183 divergence from Neanderthals and Denisovans [9]. Since huCD33 extracellular domains showed 184 high accumulation of changes compared to the great apes, we wanted to determine if these changes 185 were present in the common ancestor of the hominin lineage. We therefore compared CD33 protein 186 coding sequences from 6 Neanderthal and 2 Denisovan archaic genomes obtained from the Max 187 Planck Institute for Evolutionary Anthropology [36] (http://cdna.eva.mpg.de) with the 188 corresponding human sequences of the 1000 Genome database (Figure 1B). Interestingly, all the 189 amino acid residues in huCD33 that are different from the great apes are present in the ancient 190 genomes, suggesting their occurrence in a common ancestor. These observations thus suggest that 191 the complete loss of Sia-binding V-set domain is the latest evolutionary modification of huCD33, 192 likely succeeding the individual amino acid changes within its extracellular domain.

193 A single amino acid change facilitated CD33 engagement to the uniquely human pathogen 194 *Neisseria gonorrhoeae*. In addition to microglial expression in the brain, CD33 is also present on 195 tissue macrophages and peripheral blood monocytes [9]. These cells are important components of 196 innate immune responses throughout the body, including the reproductive tract. The human female 197 reproductive tract is also a unique niche for the microbiome, which can be invaded by important 198 pathogens like *Neisseria gonorrhoeae* (Ng). Ng is a uniquely human infectious agent, responsible 199 for the second most prevalent, sexually transmitted infection causative for the disease gonorrhea 200 in human populations. Gonorrhea affects both males and females and if untreated, can have 201 detrimental effects on reproductive health [37]. We have previously shown that Ng interacts with 202 human CD33 but not the chimpanzee ortholog [38]. The bacterium is incapable of endogenous 203 Neu5Ac synthesis, but instead scavenges the molecule from its host [39, 40]. Once inside the 204 female reproductive tract, Ng utilizes the host sugar nucleotide CMP-Neu5Ac from its 205 microenvironment to transfer Neu5Ac onto its own bacterial lipooligosaccharide. Sialylated Ng 206 then successfully interacts with several human Siglecs including 3 (CD33), 5, 9, 11, 14 and 16

[38]. However, unlike other Siglec interactions, Ng binding to CD33 appears to be entirely Siadependent. Interestingly, of all the *Neisseria* species currently known, only Ng and *Neisseria meningitidis* are pathogenic to humans and both are thought to be evolutionarily young compared to others [41]. Since reproductive health/success of an organism is the key determinant of Darwinian fitness, we hypothesized that highly infectious disease like gonorrhea could potentially impact the evolution of humans, mediated through binding immune modulating proteins like CD33.

214 To explore our hypothesis, we examined the binding of sialylated Ng to different recombinant 215 CD33 protein mutants, each containing the two extracellular domains with an amino acid residue 216 changed from human to chimpanzee at the corresponding positions identified in Figure 1B. 217 Fluorescently labelled Ng was allowed to interact with human recombinant Fc-chimeric constructs 218 of the CD33 proteins that were immobilized onto protein A-coated plates (Figure 2A). Sia-219 dependence of the interaction was confirmed by comparing binding with bacteria grown in 220 presence and absence of CMP-Neu5Ac (Supplemental Figure S1A). We observed significant 221 reduction in bacterial binding to chimpanzee CD33 (chCD33) compared to human protein 222 containing both V- and C2- domains (Figure 2B). However, in the absence of the V-set domain in 223 the truncated form of huCD33 (CD33m), bacterial binding was lost. Except for the residue at 224 position 21, all the other amino acid alterations from human to chimpanzee CD33 maintained high 225 bacterial binding. In fact, changing the amino acid residues at positions 22, 65 (of the V-set 226 domain), 152, and 154 (of C2 domain) increased the binding significantly compared to wildtype 227 huCD33. In contrast, mutating the amino acid at position 21 from human to chimpanzee residue 228 completely abolished huCD33 binding of sialylated Ng. Interestingly, mutating the chimpanzee 229 CD33 amino acid at position 21 to its corresponding human residue enabled Ng to now engage 230 chimpanzee CD33 (Figure 2C). Considering that Ng and its closest relative meningococcus are 231 both uniquely human pathogens thought to have evolved from commensal Neisseria [42], our data 232 suggest important implications of CD33 amino acid change at position 21 on Ng-huCD33 233 interaction and their mutual evolution.

Many amino acid changes in CD33 extracellular domains impact *GBS* engagement. While the association of *Neisseria* with CD33 is a case of Sia-mediated interaction, there are other examples of human pathogens that engage Siglecs in Sia-independent manner. One such example is Group B *Streptococcus* (GBS) which has been widely studied for its various ways of engaging 238 host Siglecs [43]. GBS is an encapsulated pathogen commonly associated with pneumonia, sepsis 239 and meningitis in infants and neonates. It comprises nine serologically distinct groups (Ia, Ib and 240 II -VIII), differing in their capsular sialoglycan structures, but all containing α 2-3-linked terminal 241 Neu5Ac. Certain GBS strains have been shown to bind human Siglecs 5 and 7 in a Sia-independent 242 manner through cell wall anchored β -protein [44], whereas some Sia-dependent binding was 243 observed for CD33 and Siglec-9. Human Siglec-9 binding is also thought to be partially Sia-244 independent. Interestingly, some GBS strains are also known to interact with nonhuman primate 245 Siglecs, for example, Siglec-9 from chimpanzee [25]. Since infections by GBS mostly impact 246 newborns and infants, we hypothesized that it could also play a role in overall Siglec evolution in 247 humans. Similar to the Ng-CD33 binding assay (as in Figure 2A), we examined the interactions 248 between the recombinant CD33 proteins and GBS group III strain, COHI (Figure 2D). While the 249 bacteria bound strongly with full-length extracellular domains of huCD33, the binding was 250 significantly reduced in the truncated human isoform (CD33m) and the chimpanzee protein. Like 251 Ng, GBS COHI interaction was also markedly disrupted by amino acid changes at position 21. 252 Additionally, changing the residues at positions 20 and 65 from human to chimpanzee significantly 253 reduced the bacterial interaction with CD33. However, GBS COHI engagement with the CD33 254 mutants was not entirely Sia-dependent (Figure 2E). Using GBS COHI Δ neuA, a mutant strain 255 lacking its sialyltransferase enzyme (NeuA) and hence incapable of surface sialylation, we 256 observed that about 50% of the bacterial binding to CD33 could be attributed to Sia-independent 257 interactions. Interestingly, the CD33 binding profile of COHI was not uniform for the other 258 serogroups of GBS, for example GBS group Ia strain, A909 (Figure 2F). None of the amino acid 259 changes showed significant effects on CD33 interaction with GBS A909, relative to the wildtype 260 human protein. Even the truncated human CD33 isoform (CD33m) displayed similar binding 261 suggesting that the CD33 binding for A909 is primarily Sia-independent. Unlike Ng and GBS, we 262 did not observe any differential sialoglycan binding with E. coli K1, another uniquely human 263 pathogen of newborn infants, which contains Sia polymers on its surface (Supplemental Figure 264 S1B). Altogether, the data demonstrate the diverse nature of CD33-interactions in three major 265 pathogens and suggest an impact of uniquely human pathogens in the evolution of CD33 ligand-266 binding domain.

Ancestral sialoglycan preference of CD33 is disrupted by amino acid change at position 21.
A key change in the evolution of humans was the loss of CMP-Neu5Ac hydroxylase (CMAH), the

269 enzyme that converts CMP-Neu5Ac to CMP-Neu5Gc resulting in a primarily Neu5Ac-rich 270 sialome in humans, unlike any other Old-World primates, which express both Neu5Ac and 271 Neu5Gc. This change is dated to ~2-3 mya when human ancestors were evolving from ancestral 272 hominins. Since we observed numerous changes mainly in huCD33 V-set domain which is critical 273 in sialoglycan interaction and therefore important for its downstream signaling pathways, we 274 wanted to specifically understand the effect on CD33 sialoglycan interactions. We used a 275 microarray of chemoenzymatically synthesized glycans with defined structures, terminally capped 276 with either Neu5Ac or Neu5Gc in different glycosidic linkages and examined their relative 277 interactions with recombinant, soluble CD33 proteins (Figure 3). Human CD33 with V- and C2-278 domains bound to both Neu5Ac and Neu5Gc-terminating sialoglycans and showed maximum 279 binding when the Sia was α 2-6-linked to an underlying lactose or lactosamine glycan 280 (Supplemental Figure S2). Most of this binding was lost in the truncated huCD33 lacking the Sia-281 binding V-set domain, indicating that the interactions are Sia-dependent. Conversely, the 282 chimpanzee protein (which is identical to the bonobo orthologs and differs by only two amino 283 acids from the gorilla) demonstrated strong preference towards Neu5Gc-terminating sialoglycans 284 and showed almost no binding for Neu5Ac-epitopes. Considering the varied sialoglycan profiles 285 of the two organisms, these distinct binding preferences of human and chimpanzee CD33 are 286 interesting and suggest functional implications of the evolutionary changes in their extracellular 287 domains.

288 Because the impact of amino acid residue at position 21 was most pronounced in both of our 289 bacterial-CD33 binding assays (Figure 2B and 2D), we next examined the influence of this change 290 on CD33-sialoglycan binding (Figure 3 and Supplemental Figure S2). Indeed, changing the amino 291 acid at position 21 completely altered Sia-epitope preference of CD33 for both human and 292 chimpanzee. The presence of human amino acid residue at position 21 enabled strong binding of 293 Neu5Ac-epitopes by chCD33, unlike its entirely Neu5Gc-preferring wildtype counterpart. On the 294 other hand, the chimpanzee amino acid at the same position in human CD33 abolished its Neu5Ac 295 binding. To determine if the Sia-binding changes are specific for position 21 and not an arbitrary 296 effect of any amino acid change in V-set domain, we also looked at the Sia-epitopes of position 20 297 amino acid substitutions. Unlike position 21, amino acid modifications at position 20 did not have 298 any major impact on the Sia-binding of CD33, which maintained the overall wildtype profile. 299 Interestingly, modifications at position 22 demonstrated Neu5Ac-prefered binding for huCD33,

300 while chimpanzee amino acid residues at 65 and 66 of huCD33 almost abolished any sialoglycan 301 binding. Altogether, the data emphasized the importance of different amino acid changes in 302 huCD33 V-set domain for its sialoglycan binding and identified the amino acid at position 21 to 303 be critical in the functionality of CD33 protein.

304 MD simulations provide structural insights for the differences in Sia-binding preference 305 between human and chimpanzee CD33. We performed an extensive theoretical investigation 306 based on molecular dynamics simulations. A detailed analysis of several available crystal 307 structures of huCD33 revealed that the V-set domain is dynamic. For example, the C-C' loop as 308 well as the side chains of phenylalanine at position 21 (Phe21) and histidine at position 45 (His45) 309 are resolved in two different conformations in PDB entry 5ibb (Supplemental Figure S3). Of all 310 the amino acids that differ between human and chimpanzee, only the side chain of Phe21 is in 311 direct contact with a bound Neu5Ac residue in the crystal structures of huCD33 (through the 312 methyl group at position 5). Based on the assumption that Neu5Gc binds to the same binding site 313 as Neu5Ac, the change in binding preference from Neu5Gc (in chimpanzee) to Neu5Ac (in human) 314 cannot be explained by a simple I21F mutation. Both amino acids have hydrophobic side chains 315 that cannot establish favorable interactions with the polar glycolyl group. Consequently, there is 316 probably a more complex reason for the shift of binding preference. Based on data derived from 317 47 molecular dynamics (MD) simulations covering an accumulated timescale of more than $100 \,\mu s$ 318 we conclude that in chCD33 His45 adopts mainly the 'up' conformation (Figure 4A), which allows 319 favorable hydrogen bonding with the glycolyl group. MD simulations (as well as x-ray 320 crystallography) show that in huCD33 His45 can also exist in the 'up' conformation (Figures 4C 321 and Supplemental Figure S4), which would be compatible with favorable Neu5Gc binding. 322 However, when His45 is in the 'down' conformation Phe21 can stack partly with tyrosine (Tyr) at 323 position 127 (Figure 4B) forming a small hydrophobic pocket, which allows the methyl group of 324 Neu5Ac to bind favorably. To demonstrate if the binding affinity difference between Neu5Ac and 325 Neu5Gc may be indeed correlated to the up/down conformational equilibrium of His45, we 326 performed a series of MD simulations of chCD33 on the microsecond timescale where 327 Neu5AcOMe or Neu5GcOMe molecules are present in the solution. The lifetimes of the 328 complexes spontaneously formed during the MD with Neu5Gc are on average much longer when 329 His45 is 'up' (Figure 4D top, Supplemental Figure S4). In contrast the lifetimes of the complexes 330 spontaneously formed with Neu5Ac are much shorter independent of the conformational state of

His45 (Figure 4D bottom), which would explain the lack of measurable binding affinity of Neu5Ac
to chCD33. In summary, our extensive MD simulations - including unbiased simulation of
carbohydrate binding and unbinding events - could provide a reasonable explanation for a change
in binding specificity that is likely to be caused by an alteration of the protein-ligand interaction
pattern remote from the mutated amino acid.

336 Human-specific polymorphisms in cognitive-health related genomic variants are present in 337 all human populations. In an earlier study we observed several genes, directly associated with 338 neurodegenerative diseases or correlated with aggravation of the cognitive decline in aged-339 population, are derived alleles in humans [9]. Increasing evidence of correlation between cognitive 340 health and non-neurological, metabolic conditions, e.g., diabetes [45, 46] suggest that such derived 341 alleles could be important in the maintenance of cognitive health in human grandparents. Here, we 342 expanded this list of cognition-protective gene variants through literature and database (https://alzoforum.org) searches [20, 47-58] to include additional gene variants, namely BINI, 343 344 ARID5B, PICALM, PILRA. Supplemental Table S1 describes the characteristics of 13 human 345 genes that are implicated in diseases including dementia, cardiovascular diseases (CVD), hypertension and AD. While some of these physiological abnormalities like salt retention, 346 347 hypertension, diabetes, appear non-neurological, they have been associated with the aggravation 348 of the pathologies resulting in late-life cognitive decline [59]. Notably, the derived alleles are 349 common and found in globally diverse human populations, indicating that they predate the 350 common ancestor of modern humans (Supplemental Table S2).

351 SNPs associated with human-specific cognitive protective alleles are unusual in their absence 352 in the archaic hominin genomes. With the availability of genomes from extinct archaic hominins 353 [36, 60, 61], a set of SNPs can be assessed as to whether their protective phenotypes arose recently 354 in the evolutionary history of anatomically modern humans. We previously showed many other 355 human-chimpanzee differences were shared with archaic hominins (Denisovan/Neanderthal) 356 genomes; for example, genomic changes in CD33rSiglecs [62]. To gain similar insights about the 357 evolutionary origin of these cognitive-protective loci, we analyzed the Neanderthal and Denisovan 358 reference genomes and compared them with modern human sequences. Analysis of the 1000 359 Genomes dataset shows the presence of protective alleles in human populations with variable 360 frequency (Table 1). Analysis of the available genomic data from Neanderthal and Denisovan 361 genomes showed that only two derived variants (rs2975760 and rs2588969; Table 1) are present

362 in these archaic genomes, suggesting the remaining eleven derived, protective variants arose after 363 the divergence of modern and archaic hominins approximately 0.5 mya [35, 63]. This is in striking 364 contrast to most human-chimpanzee genomic differences in which the archaic hominins are similar 365 to humans. In fact, majority of the Sia-related genes lack positive selection signatures and rather 366 show neutral evolution in the modern human lineage [64]. To more formally assess whether the 367 high frequency, global distribution, and recent origin observed for eleven of the thirteen SNPs is 368 unusual, we performed a resampling analysis of variants in the genome. Variants in the 1000 369 genomes dataset that met the following criteria were considered: 1) present in both Altai 370 Neanderthal and Denisovan minimal filters, 2) derived in at least one modern individual from non-371 admixed African populations, 3) called in both archaic samples, and 4) have an ancestral allele 372 matching the reference or alternative allele. To eliminate any bias in the analysis and match the 373 allele frequency (AF) of these SNPs compared with that of any random SNPs, we first matched 374 our universe of SNPs to the 13 SNPs of interest by AF, ± 2 derived haplotypes (Figure 5). 375 Resampling was then performed by drawing a SNP from each of the 13 matched sets and assessing 376 how many derived alleles were observed, resulting in a p-value = 0.08333 ± 0.00003 . As a less 377 conservative estimate, directly sampling from the universe of SNPs and estimating the probability 378 of observing at most two derived SNPs and a mean allele frequency as large as the empirical 379 variants of interest produced a highly significant p-value = 0.00487 (Supplemental Figure S5). 380 Repeating either analysis on the set of other Siglec-related SNPs indicates they are consistent with 381 a random draw from the genome [62]. Regardless of the individual limitations, taken together our 382 phylogenetic analyses demonstrate the unique patterns of allele frequencies in worldwide 383 populations distribution of these thirteen late-life cognitive decline linked SNPs (Figure 5 and 384 Supplemental Figure S5). Interestingly, co-inherited CD33 SNPs associated with the cognitive 385 health in LOAD are present only in modern human genomes [9]. A noteworthy example in our list 386 is the human gene encoding the protein, apolipoprotein E (APOE), involved in fat metabolism in 387 mammals. APOE gene exists in three allelic variants (E2, E3 and E4) where APOE4 is associated 388 with high risk of LOAD and other allele like APOE2 is protective against the cognitive decline in 389 elderly caregivers [65]. Interestingly the presence of APOE4 is also correlated with the protection 390 from severe diarrhea in children [66]. While conclusive determination of the positive selection of 391 these alleles in modern human requires further analysis, our data suggest that the evolutionary 392 origin of most of these cognitive-health protective changes followed the divergence of modern

393 humans from archaic genomes. This is also supported by the presence of grandparents, uniquely

in humans. Regardless, the process of evolutionary emergence of each of these alleles is likely to

395 be distinct and deserves further investigation.

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Discussion

398 Fossil evidence and genomic comparisons leave little doubt about the fact that our species evolved 399 from an African hominin. However, a detailed understanding of modern human origins is plagued 400 by numerous uncertainties, with regard to the identity of the ancestral lineage and precise 401 geographic locations. Evolution of modern humans was accompanied by many anatomical and 402 behavioral changes, but increasing evidence suggests it also included uniquely human-derived 403 modifications in the genome compared to the archaic genomes (Neanderthal/Denisovan) or the 404 genome of the great apes [9, 62]. Taken together with our previous study [9], we have identified 405 many such human-specific genes associated with cognitive health of grandmothers and other 406 human elders who are often involved in the caregiving of the young. These findings, which appear 407 paradoxical to the concept of senescence due to antagonistic pleiotropy, have lent much additional 408 support to the "Grandmother hypothesis" [1] bolstering the case for selection of human female 409 post-reproductive survival and the existence of grandmothers. Unlike in any other mammals 410 (except orcas and some other toothed whales), the occurrence of this prolonged post-reproductive 411 life span in humans has stirred scientific interest. While deciphering the precise evolutionary 412 course of any gene/protein is challenging and the proposed schemes/players are not entirely 413 verifiable, here we attempt to compile the current evolutionary and experimental findings of one 414 such protein associated with late-life cognitive-decline: CD33.

415 A ratio of high wildtype human CD33 and a low truncated isoform of CD33 have been implicated 416 in the progression of LOAD associated with the cognitive health of elderly population. In contrast, 417 LOAD is unknown in chimpanzees, although evidence of LOAD pathologies has been observed 418 in some chimpanzee brains. We found that human CD33, which is highly expressed in microglia 419 of the human but not chimpanzee brain, recognizes Neu5Ac – the predominant Sia synthesized in 420 humans – as self-associated molecular patterns (SAMPs). In contrast, our closest evolutionary 421 relative, the apes and other Old-World primates contain both Neu5Ac and Neu5Gc. We found that 422 the ancestral form of CD33 in chimpanzees and other great apes selectively recognizes only 423 Neu5Gc-glycans as SAMPs (Figure 3). Notably, Neu5Gc – the ligand recognized by chCD33 is

424 rare in chimpanzee brain, and there is also significantly less chCD33 protein compared to CD33 425 in humans [9]. On the other hand, SNPs resulting in the truncated CD33 have only been observed 426 in the human genome and not any of the archaic or great ape genomes. We also find that the 427 truncated human CD33 does not interact with Sia (Figure 3). Taken together, these observations 428 suggest that full-length CD33-Sia interactions are stronger in human brain compared to 429 chimpanzee and the human-specific SNPs in CD33 resulting in the truncated protein abolish this 430 interaction. The question remains what could have possibly led to the selection of the truncated 431 isoform of human CD33 that does not interact with Sia. In this regard, CD33 on macrophages 432 plays crucial roles in different immune responses as well as during infections. Human CD33 has 433 also recently been shown to be involved in immunomodulation during infection with hepatitis B 434 virus [29]. Our previous and current data show that uniquely human pathogens like *Neisseria* and 435 GBS display Neu5Ac that is recognized as 'self' by human but not chimp CD33 [38]. In the current 436 work, we further found that the Sia-binding-domain-depleted, truncated human CD33 isoform 437 doesn't bind and thus escape exploitation by sialylated pathogens (Figure 2). This suggests that 438 this truncated CD33 may have been an adaptation to counter the CD33-exploiting, immune-439 evasive behavior of pathogens like Neisseria and GBS.

440 Taking together all currently available experimental data (including this study) we attempt to draw 441 a plausible evolutionary scenario for CD33 protein evolution in humans and present in the context 442 of relevant evolutionary events (Figure 6). We hypothesize that the scarcity of the strongly 443 preferred Neu5Gc ligand of ancestral CD33 in the brains of chimpanzee (and other great apes) was 444 associated with low microglial expression. Subsequent hominin loss of CMAH (i.e., complete loss 445 of Neu5Gc ligand) could then have selected for the upregulation of CD33 levels perhaps to 446 compensate for the loss of ligands, a change to Neu5Ac-binding preference, and functional 447 recruitment of CD33 to human microglia. Alongside the microglial CD33, the corresponding 448 changes in the tissue macrophage proteins might have facilitated the emergence of Neu5Ac-coated 449 pathogens (for example, N. gonorrhoeae and Group B Streptococcus) that evolved "molecular 450 mimicry" of Neu5Ac-SAMP ligands to manipulate the immune response. Appearance of the 451 truncated isoform lacking the ligand-binding domain (CD33m), then probably allowed CD33 to 452 escape the immune evasion by these sialylated pathogens (Figure 2). This selection pressure to 453 stop manipulation by sialylated pathogens could have also altered splicing towards a higher level 454 of truncated CD33, which also gets diverted to peroxisomes [12]. While the significance of this

455 diversion is unclear, decrease of full-length CD33 would facilitate escape from Neu5Ac-coated, 456 CD33-engaging pathogens. Finally, sometime during the last 1 million years, increased brain size 457 presumably selected for early, short interbirth interval in human, which might have resulted in 458 more helpless young, requiring cooperative breeding and caregiving. However, the value of 459 postmenopausal grandmothers and other elderly caregivers would then have been blunted by the 460 appearance of LOAD. The synthesis of the truncated isoform of CD33 protects from *Neisseria* 461 during reproductive age and a higher ratio of truncated to full-length isoforms correlates to decrease of LOAD in grandmothers. However, a small amount of the full-length isoform remains, 462 463 likely to downregulate hyper-inflammation that might arise during prolonged absence of SAMP-464 recognition. Notably when an elderly caregiver gets LOAD, not only are the evolutionary benefits 465 of the individual lost, but this also presents an increased burden to care for that elder individual. 466 Altogether under this proposed scenario, the current state in the evolution of human CD33 protein 467 represents a trade-off between the evolutionary response to exploitation by pathogens in early life 468 and cognitive maintenance in post-reproductive late life.

469 A similar evolutionary scenario appears to underly the case of the human APOE gene 470 where variants include both risk alleles (APOE4) and protective alleles, (APOE2, and APOE3) for 471 CVD and LOAD [65]. In this instance, the ancestral APOE4 allele is associated with increased 472 risks of loss of cognitive functions and the derived alleles may serve to protect the cognition of the 473 elderly caregivers. Interestingly the APOE4 allele is also correlated with the protection from severe 474 diarrhea in early years of life [66]. Given these examples like APOE and CD33, it remains to be 475 seen how widespread this evolutionary pattern is wherein variants conveying survival advantages 476 in early life coexist with other variants that protect cognition late in life.

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483

484 Experimentation (S.S., N.K., T.C., A.V., A.S., S.D., M.F.), Data analysis (S.S., N.K., T.C.,
485 A.V., A.S., S.D., J.M.A., M. F., P. G., A.V.), Critical reagents (H.Y., X.C.), Original draft (S.S.,

Author Contribution

Declaration of interest

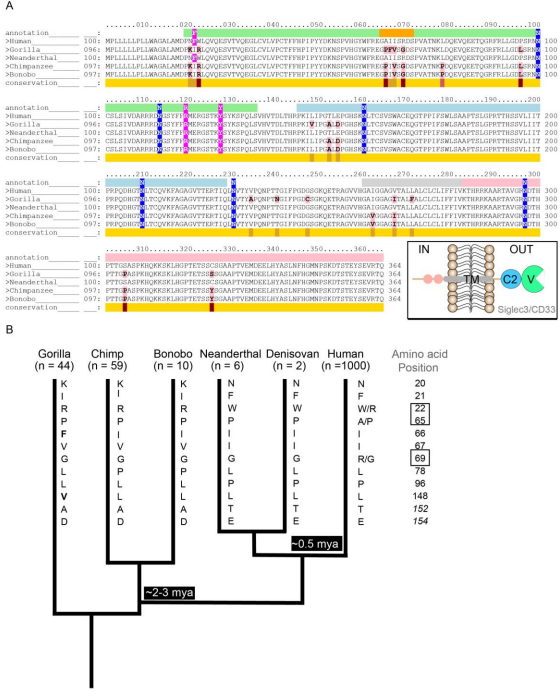
- 486 P.G., A.V.), Writing (S.S., N.K., T.C., A.V., A.S., X.C., J.M.A., M.F., P.G., A.V.), Overall
- 487 supervision (J.M.A., P.G., A.V.), Funding acquisition (A.V.).
- 488 Data Availability
- 489 The data for the resampling analysis is available at Code Ocean.
- 490
- 491
- 492 The authors have declared that no conflict of interest exists.

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Figures





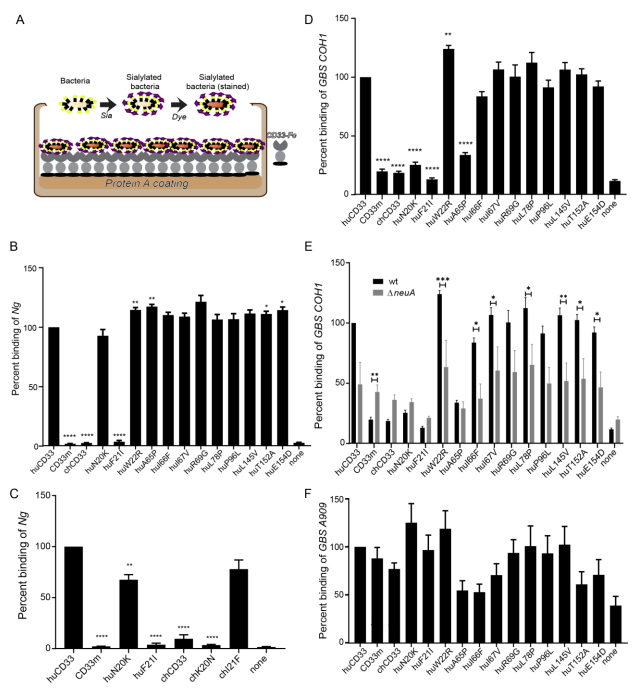
494

495 Figure 1: Human specific changes in CD33 are primarily present in the Sia binding V-set domain. (A) Comparison of amino acid sequences of CD33 from humans and "great apes" was 496

497 performed using Conformational Analysis Tools software. The Great ape genomes included in the

498 analysis are gorilla, chimpanzee and bonobo and were compared against the human protein as the

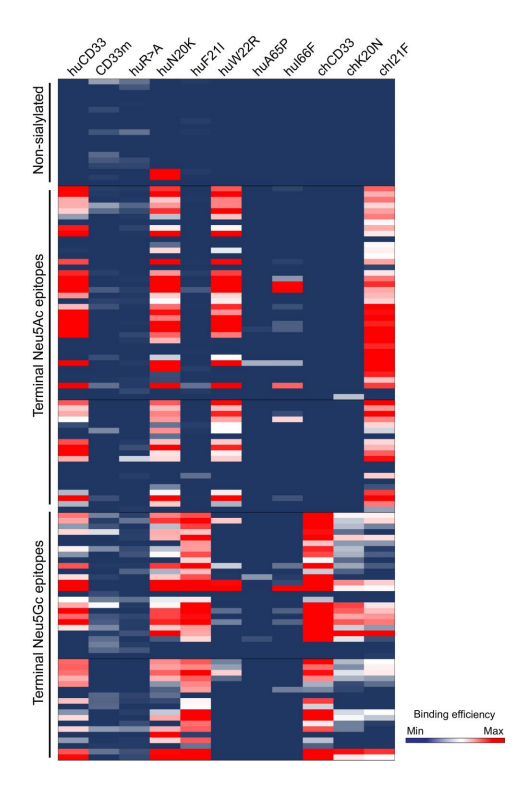
499 template. The conservation of the sequence is indicated with yellow being the most and red being 500 the least conserved regions. Amino acids that are different from huCD33 are highlighted in pink. 501 Amino acids encoding the different CD33 domains are indicated above the sequence with different 502 colors corresponding to schematic in inset, namely, V-set domain in green, C2 domain in light 503 blue, transmembrane domain in grey and cytoplasmic end in light pink. The flexible C-C' loop is 504 indicated in orange. Amino acids that are in contact with Neu5Ac in huCD33 are highlighted in 505 magenta and N-glycosylation sites in blue. (B) Phylogenetic analysis of the evolution of the 506 extracellular domains of CD33 proteins from human, Great apes and archaic genomes. The number 507 of genomes (n) for each group included in the analysis is indicated. Human and great ape CD33 508 sequences were compared with six Neanderthal and two Denisovan genomes. Amino acid changes 509 present in human CD33 were also present in the ancient genomes. The positions of the amino acids 510 that are different between human and the apes are mentioned, and the identity of the amino acid 511 present in the corresponding positions for each group is indicated by the single letter abbreviations 512 along the branch. Amino acids at positions 152 and 154 are within the C2 domain of CD33 protein 513 and italicized. Polymorphisms within the human population at positions 22, 65 and 69 of CD33 514 protein are indicated. Amino acids in gorilla CD33 at positions 66 and 148 are different from other 515 apes and are bold. Possible timeline for the diversion of the hominin lineage is indicated in the 516 tree. Length of the branches in the tree is not to scale. Mya = million years ago.



517

Figure 2: Human specific amino acid changes in CD33 affects bacterial binding. (A) Schematic of the ELISA-based assay using recombinant CD33-Fc chimeric proteins immobilized on protein A coated plates used to determine binding of the sialylated bacteria is shown. (B) Binding of fluorescently labelled *Neisseria gonorrhoeae* (*Ng*) was determined. The position of the amino acid different from the wildtype human CD33 protein is indicated below each bar in the xaxis. The bacterial binding to each individual CD33 mutant was normalized to the binding of

524 wildtype human CD33 for that assay. "None" indicates no protein control for the background 525 bacterial binding to the plate. (B) Binding of *Neisseria* to immobilized recombinant CD33 proteins 526 containing the corresponding amino acid mutation (position 20 or 21) in either in human or chimp 527 CD33 protein backbone. (D) Binding of Group B Streptococcus (GBS) COH1 strain to different 528 CD33 mutant proteins in an ELISA based assay with immobilized recombinant CD33 proteins. 529 (E) Sialic acid dependence of the binding was determined using wildtype and $\Delta neuA$ mutant strains 530 of COH1. (F) Interaction of CD33 proteins among different GBS strains was compared using A909 531 and COH1 strains. 'hu' indicates the corresponding amino acid change in human CD33 backbone 532 and 'ch' using chimp CD33. The graphs show the cumulative result from 3 independent 533 experiments, each done in triplicate. Statistical analysis was performed in Prism software using one-way ANOVA with Durrett post comparison test. * < 0.01, ** < 0.001, *** < 0.0001. 534



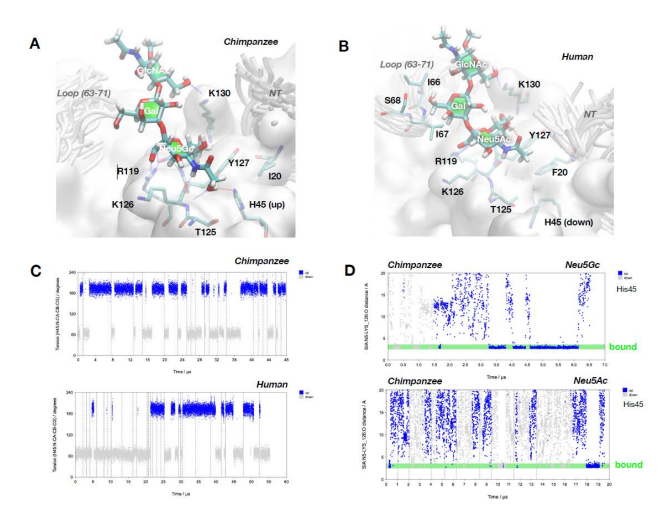


536

Figure 3: Single amino acid changes affect CD33 sialoglycan binding. Sialoglycan binding
 profile of purified, soluble, recombinant CD33 proteins was determined using a sialoglycan
 microarray containing defined, chemically synthesized glycans. Non-sialylated, Neu5Ac- and

540 Neu5Gc-terminating glycans were grouped together in the heatmap as shown in the left. Each

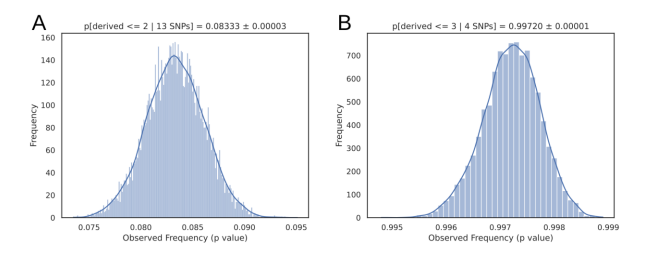
- 541 column indicates the binding profile of the protein indicated on the top and each row represent a
- 542 distinct glycan. Blue indicates no binding and red indicates very strong binding preferences
- 543 characterized by an average relative fluorescence unit (RFU) of more than 90th percentile. The
- result of the heatmap is summarized in Supplemental Figure S2 and the names of the individual
- 545 glycans are presented in Supplemental File S1.



546

547 Figure 4: Structural modeling to understand the differential binding preference of human 548 and chimpanzee CD33 proteins. (A) 3D model of the complex between Neu5Gca2-3GalB1-549 4GlcNAcβOMe and chCD33. The increased affinity of Neu5Gc may be explained by 550 intermolecular hydrogen bonds involving the OH-group of Gc. It should be noted that the number 551 of favorable interactions is maximal when His45 is in 'up' conformation. (B) 3D model of the 552 complex between Neu5Ac α 2-3Gal β 1-4GlcNAc β OMe and huCD33. The methyl group of Ac is 553 located in a small hydrophobic pocket formed by the side chains of Tyr127 and Phe20. It should 554 be noted that His45 is in 'down' conformation because otherwise - in the conformation shown -555 the bulky side chain of Phe20 would overlap partly with His45 in 'up' conformation. (C) Molecular 556 dynamics of His45 side chain orientation. Accumulated MD trajectories of torsion angle N-Ca- $C\beta$ - $C\gamma$ are shown. The 'up' conformation is present when torsion values are fluctuating around 557 558 200 degrees and the 'down' conformation is characterized by values around 70 degrees. For 559 chimpanzee, it can be observed reproducibly that simulations started with His45 in 'down'

560 conformation undergo a transition to the 'up' conformation on the microsecond timescale. In 561 contrast the 'down' conformation appears to be more stable in huCD33, which would make 562 Neu5Ac binding more likely. (D) MD simulation of unbiased binding and unbinding events of 563 Neu5Gc (top) and Neu5Ac (bottom) to chCD33. For Neu5Gc the lifetime of the complex is 564 significantly longer when His45 is in 'up' conformation, as can be seen from the 6.5 µs MD 565 simulation shown on the top. Also, for Neu5Ac multiple binding and unbinding events occurred 566 on a timescale of about 20 µs, however in general (with one exception) the lifetimes of the 567 complexes formed are significantly shorter than for Neu5Gc.



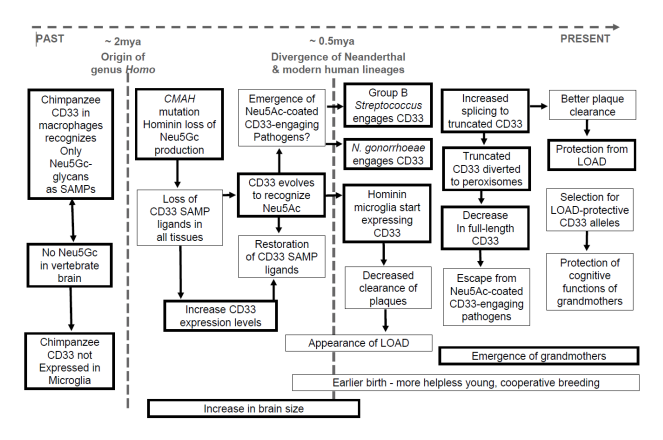


569 Figure 5: Resampling analysis with matched allele frequency SNPs from 1000 genomes

570 variants. Frequency distribution of SNPs with similar properties to the LOAD protective set (A)

571 and other Siglec SNPs (B).

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572

573 Figure 6: Scenario for evolution of human CD33 in relationship to cell surface sialic acids, 574 infectious disease, brain microglia and cognitive maintenance of grandmothers and other 575 elderly caregivers. This schematic presentation combines the known/likely facts (thick-outlined 576 boxes, including data from this manuscript) as well as suggested possibilities (thin-outlined boxes) 577 into the most likely evolutionary scenario for human-specific evolution of CD33. Starting from 578 the left, the likely chronological order of occurrence is indicated (by arrowheads) with the 579 approximate timeline on the top, along the dotted lines. '?' indicates our reasonable assumption 580 leading to the event. See text for further discussion.

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581 582

Table

583 Table 1: Gene variants directly or indirectly affecting cognitive function. Allele has the derived

584 allele as the lower, bolded entry. Archaic genotypes are reported for SNPs passing all quality

585 filters.

				Archaic Genotype				
Gene	Associated Disease ^a	SNP ID	hg19 Position	Allele	Derived Global AF ^b	Altai	Denisovan	
AGT	Sodium retention	rs699	1:230845794	G A	0.295	0/0	0/0	
BIN1	AD	rs7561528	2:127889637	A G	0.8	0/0	0/0	
SGC2	Hypertension	rs1017448	2:224466344	T C	0.879	0/0	0/0	
CAPN10	Type II Diabetes	rs2975760	2:241531163	C T	0.882	1/1	1/1	
PPARG	Type II Diabetes	rs1801282	3:12393125	C G	0.070	0/0	0/0	
CYP3A5	Sodium retention	rs776746	7:99270539	T C	0.621	0/0	0/0	
ARID5B	AD	rs2588969	10:63611354	A C	0.532	0/0	1/1	
SPON1	Dementia	rs2618516	11:14021639	C T	0.341	0/0	0/0	
PICALM	AD	rs10792832	11:85867875	G A	0.313	0/0	0/0	
FICALM		rs3851179	11:85868640	C T	0.315	0/0	0/0	
APOE	LOAD	rs429358	19:45411941	C T	0.849	0/0	0/0	
AIOL		rs7412	19:45412079	C T	0.075	0/0	0/0	
CD33	LOAD	rs3865444	19:51727962	C A	0.211	0/0	0/0	
PILRA	AD	rs1859788	7:99971834	G A	G 0.341		-	
TCFLC2	Type II Diabetes	rs7903146	10:114758349	T C	0.772	-	-	
CD33	LOAD	rs12459419	19:51728477	C T	0.211	-	-	

586 ^a See supplemental Table S1 for details and the primary literature citations.

587 ^b See supplemental Table S2 for the global population distribution.

588	Methods
589	Bacterial culture and cell lines. The bacterial strains used were <i>Neisseria gonorrhoeae</i> $F62\Delta lgtD$
590	(generous gift from Sanjay Ram, University of Massachusetts Worcester), Group B Streptococcus
591	(GBS) strains COH1wt, COH1ΔneuA, A909wt and A909ΔneuA (generous gifts from Victor Nizet,
592	University of California San Diego). Neisseria were grown overnight on chocolate II agar plate
593	and GBS on Todd Hewitt agar plate at 37 $^{\circ}\!C$ and 5% CO ₂ from the respective frozen glycerol
594	stocks. Prior to the assay, GBS was grown in Todd Hewitt broth at 37 $^\circ$ C and 5 $^\circ$ CO ₂ without
595	shaking. The E. coli K1 strain was grown in LB. For the CD33 protein purification, HEK293A
596	cells were grown in DMEM media (Invitrogen) containing 10% FCS at 37 °C and 5 % CO ₂ .
597	Sialylation of Neisseria. Following overnight growth on chocolate agar plate, the bacteria were
598	grown in GC broth supplemented with IsoVitaleX at 37 °C, 5% CO2 and shaking at 200 rpm in
599	presence or absence of 30 µM CMP-Neu5Ac (Nacalai USA. Inc.) until OD600 equivalent to 0.4-
600	0.5.
601	Bacterial staining. Following appropriate growth, the bacteria were washed with pre-warmed
602	HBSS and stained with 2 μM SYTO13 (Thermo Scientific) for 30 min at 37 °C and shaking at 200
603	rpm in dark. After incubation, the stained bacteria were washed with HBSS and resuspended to a
604	final concentration of $OD600 = 1/ml$ in HBSS for the binding assay.
605	Generation of CD33 mutant proteins. A genomic fragment (1228 bp) of human or Chimpanzee
606	CD33(M), including the first 4 exons (2 Ig domains) was fused with pcDNA3.1(-) containing a C-
607	terminal FLAG (EK) sequence followed by a hIgG1-Fc genomic fragment (hinge + 2 Ig-like
608	domains) and described elsewhere [67, 68]. Sixteen mutant variants were made from either
609	construct above using New England Biolabs Q5 site directed mutagenesis Kit according to the
610	manufacturer's instructions (Supplemental Table 3). Mutagenesis primers listed were designed
611	using NEBaseChanger software.
612	Truncated CD33(CD33m) _EK_Fc Construction: U937 cells were cultured in RPMI 1640
613	supplemented with 10% FCS. Total mRNA was isolated using Qiagens Oligotex Direct mRNA
614	Mini Kit according to the manufacturer's instructions. CD33m was amplified by PCR using
615	SuperScript III One-Step RT-PCR (Invitrogen) and Gene-specific primers 5'-
616	TTATATGCTAGCGCCACCATGCCGCTGCTGCTGCTGCTGC-3', NheI site underlined and
617	5'-GCGCGCGCGATATCATGAACCACTCCTGCTCTGGTCTCTTG-3', EcoRV site underlined.
618	PCR products were run on 2% agarose gel and the 396 bp bands corresponding to CD33(m) were

excised and cut with NheI/EcoRV restriction enzymes. Digested bands were sub-cloned into
pcDNA3.1(-) containing a C-terminal FLAG (EK) sequence followed by a hIgG1-Fc genomic
fragment (hinge + 2 Ig-like domains).

622 **Purification of CD33 mutants.** Transfection supernatants were collected and spun down at 500 623 g for 5 mins to remove cellular debris. The pH of each supernatant was adjusted to pH 8.0 for 624 optimal binding of protein A-Sepharose beads to hIgG Fc fusion protein. Protein A-Sepharose 4 625 Fast Flow suspension (GE Healthcare) was washed with Tris-Buffered Saline (TBS) pH 8.0, and 626 a 1:500 ratio of beads: media added to each supernatant. Each tube was subsequently incubated for 627 24 hrs on a roller in the cold-room. After 24 hours supernatants plus beads were transferred to 628 disposable columns until all liquid has run thru. Beads were washed 3x with TBS pH 8.0 before 629 being eluted directly in 0.3 ml of 1 M Tris-HCl pH 8.0 using 0.1 M Glycine Buffer pH 2.8. Each 630 eluate was put into an Amicon Ultra-15 filter unit with MWCO 30 K for each full length CD33-631 EK_Fc variant and MWCO 10 K for huCD33m-EK_Fc. Tubes were centrifuged at 4,000 g for 20 632 mins. Run-through was discarded and the columns washed 3x with TBS pH 8.0. After the last 633 wash, each retentate was recovered from the column and stored at -80°C.

634 Binding assay with the bacteria. Bacterial binding with the CD33 proteins were done with the recombinant Fc-chimeric proteins of CD33. Briefly, protein A coated black 96-well plate (Pierce, 635 636 Thermo Scientific) was washed thrice with TBS containing 0.05% Tween 20 (TBS-T) and coated 637 with 200 ng/well of the respective CD33 protein diluted in 200 mM Tris-HCl pH 8.0, 150 mM 638 NaCl and 1% BSA at 4 °C overnight. Following incubation, the coated plate was washed with 200 639 mM Tris pH 8.0, 150 mM NaCl to eliminate the unbound proteins. Stained bacteria equivalent to 640 OD600 = 0.1 was added to each well of the plate and allowed to interact with the proteins for 30 641 min at 37 °C and 5% CO₂ without shaking. Following incubation, the plate was washed with TBS-642 T to eliminate any unbound bacteria and the residual fluorescence was measured upon excitation 643 at 488 nm and emission at 530 nm. The data were analyzed using the excel and Prism software. 644 Evolutionary analysis and Detection of positive selection. The protein coding sequences of 645 CD33 were aligned using CLUSTAL W program implemented in MEGA7 and then back 646 translated to obtain a codon alignment. The phylogenetic tree of CD33 protein coding sequences

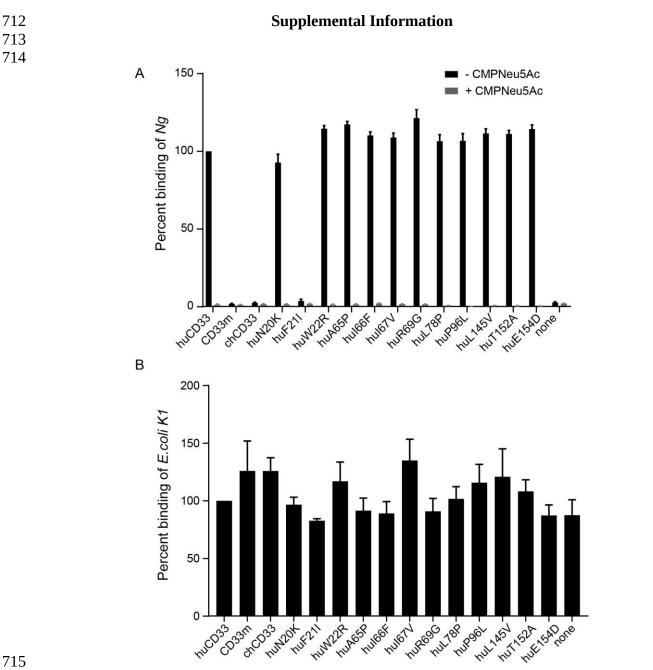
were reconstructed with neighbor-joining method which was implemented in MEGA7 (Figure 1),
1000 bootstrap replicates [69]. The unrooted neighbor joining tree was used for the subsequent
analysis.

650 VCF files were accessed from ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/ for 651 1000 genomes project, http://cdna.eva.mpg.de/neandertal/altai/AltaiNeandertal/VCF/ for Altai 652 Neanderthal and http://cdna.eva.mpg.de/denisova/VCF/hg19_1000g/ for Denisovan. Quality 653 filters were obtained from https://bioinf.eva.mpg.de/altai_minimal_filters/ for Altai and 654 Denisovan. Individuals in 1000 genomes datasets were assigned to populations using 655 http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/working/20130606 sample info/20130606 s 656 ample_info.txt. First, all vcf files were filtered by intersecting with the quality bed files using 657 bedtools intersect (v2.28.0). The filtered vcf files were then combined, per chromosome, to match 658 their position, reference and alternative allele using a custom python script. VCF information was 659 retained along with per-population allele frequencies and archaic genotypes. Next, ancestral alleles 660 were obtained from ensembl (https://rest.ensembl.org/variation/homo sapiens) by querying each 661 SNP id and appending to the joint vcf entries. The joint vcf files were used as input for further 662 processing in a jupyter notebook to perform resampling analysis. Each SNP of interest is used to 663 select collections of SNPs with matching global allele frequencies, +/- 0.004, or +/- 2 observed 664 haplotypes. A single draw consists of selecting one SNP from each collection to produce a 665 simulated observation and the number of SNPS with derived archaic haplotypes are recorded. 666 After 10,000 such draws, the faction of draws with fewer or equal numbers of derived SNPs is 667 used to produce a p-value estimate. The process is repeated 10,000 times to produce a histogram 668 and provide a confidence estimate on the reported p values (+/- SEM). Methods to replicate the 669 analysis can be found on Code Ocean.

670 Non-synonymous/ synonymous substitution ratios ($\omega = dN/dS$, or Ka/Ks) have become a useful 671 means for quantifying the impact of natural selection on molecular evolution. In general, the ratio 672 $\omega = dN/dS$ is less than one if the gene is undergoing purifying selection, equal to one if the gene 673 is evolving neutrally, and greater than one if positive selection has accelerated the fixation of non-674 synonymous substitutions that resulted in amino acid changes. The pair-wise computation of 675 Ka/Ks between V-set exon of each species were performed using the program DnaSp v.0 6.0. The 676 initial unrooted tree fed to the program in the format of Newick was: ((Chimpanzee5:0.00000000, 677 Bonobo:0.0000000):0.00222522, Gorilla:0.00979959, Human:0.01351877).

Molecular Simulation. Starting structures of the V-type domain (residues 18-142) of CD33 were built based on PDB entries 5j0b (A chain) and 6d49 using the graphical interface of YASARA [70]. The two structures differ significantly with respect to the conformation of the C-C' loop 681 (residues 63-71, compare Supplemental Figure S3). A single mutation (G69R) was introduced into 682 5j0b to build CD33(human). The initial 3D models of chCD33 were built by swapping residues: 683 N20K, F21I, W22R, A65P, I67V, R69G (in 6d49), L78P, P96L. An N-glycan core (M3) was 684 attached to Asn100. The side chain of His45 was modeled in two conformations (compare Figure 685 6): 'down' (as in PDB entry 6d49) and 'up' (as present in PDB entries 5ihb or 5j06 chains A). The 686 systems were solvated in 0.9% NaCl solution (0.15 M) and simulations were performed at 310 K 687 using periodic boundary conditions. The box size was rescaled dynamically to maintain a water 688 density of 0.996 g/ml. Additionally systems were built that contain five molecules of 689 Neu5GcaOMe or Neu5AcaOMe distributed in the simulation box which allowed to simulate 690 binding events. Simulations were performed using YASARA with GPU acceleration [71]. In total 691 27 MD trajectories were sampled for huCD33 and 20 for chCD33, most of them covering a 692 microsecond timescale (compare Supplemental Figure S4). Conformational Analysis Tools (CAT, 693 http://www.md-simulations.de/CAT/) was used for analysis of trajectory data, general data 694 processing and generation of scientific plots. VMD [72] was used to generate molecular graphics. 695 **Sialoglycan microarray**. The sialoglycan microarray experimental method was adopted from the 696 literature reported earlier [73, 74]. Chemoenzymatically synthesized sialoglycans were quantitated 697 utilizing DMB-HPLC method [75] and 10 mM aqueous stock solutions were prepared. Next, the 698 glycans were diluted to 100 μ M in 300 mM Na-phosphate buffer (pH 8.4) and printed in 699 quadruplets on NHS-functionalized glass slides (PolyAn 3D-NHS; catalog# PO-10400401) using 700 an ArrayIt SpotBot® Extreme instrument. The slides were blocked using 0.05M ethanolamine 701 solution in 0.1 M Tris-HCl (pH 9.0), washed with warm Milli-Q water and dried. Printed slides 702 were fitted in a multi-well microarray hybridization cassette (ArrayIt, CA) and rehydrated using 703 400 µl of ovalbumin (1% w/v, PBS) for one hour in a humid chamber with gentle shaking. The 704 solution was discarded followed by the addition of 400 µl solution of the CD33 protein (30 µg/ml 705 in PBS with 1% w/v ovalbumin) in the individual well. The slides were incubated for 2 h at ambient 706 temperature with gentle shaking followed by washing with PBS-Tween (0.1% v/v) and PBS. The 707 wells were then treated with Cy3-conjugated goat anti-human IgG (1:500 dilution in PBS), 708 incubated for 1h in a dark humid chamber with gentle shaking. After washing and drying, the slides 709 were scanned using a Genepix 4000B scanner (Molecular Devices Corp., Union City, CA) at 710 wavelength 532 nm. Data analysis was performed using the Genepix Pro 7.3 software (Molecular 711 Devices Corp., Union City, CA).

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717 Supplemental Figure S1: While E. coli does not bind CD33, human CD33 binding by 718 Neisseria is Sia-dependent. (A) Binding of Ng with wildtype or mutant CD33 proteins was 719 determined in the same manner as in Figure 2A. The bacteria for the assay were either grown in 720 presence (+) or absence (-) of exogenous CMP-Neu5Ac as indicated in the legend. All the binding was normalized to wildtype human CD33 binding. Cumulative data from 2 independent 721 722 experiments, each done in triplet is presented. (B) Binding of E. coli K1 was determined using the

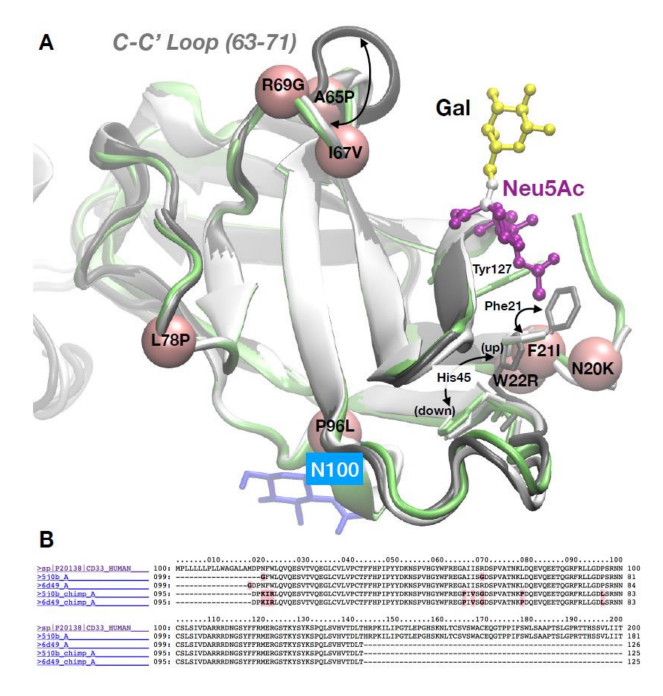
- 723 different CD33 proteins. None of the proteins showed any increased binding to the bacteria relative
- to no protein (control) containing blank well, indicating that there is no binding of the bacteria
- with the protein.

Sialic acid	Linkage	huCD33	CD33m	huR>A	huN20K	huF211	huW22R	huA65P	hul66F	chCD33	chK20N	chl21F
Non-Sia		-	-	-	_*	-	-	-	-	-	-	-
	α2-3	++	-	-	++	-	+++	-	_*	-	-	++
Neu5Ac	α2-6	+++	-	-	+++	-	+++	-	-	-	-	+++
	α2-8	-°	-	-	+	-	-°	-	-	-	-	+++
Neu4,5Ac ₂	α2-3	+	-	-	+	-	+	-	-	-	-	+++
	α2-3	++	-	-	-/+	-	+	-	-	-	-	+
Neu5,9Ac ₂	α2-6	-/+	-	-	+	-	-/+	-	-	-	-	+++
Neu5Ac8Me		-	-	-	-	-	-	-	-	-	-	-
	α2-3	+	-	-	++	+++	_ °	-	-*	+++	-/+	-
Neu5Gc	α2-6	+++	-	-	+++	+++	-	-	-	+++	+++	++
	α2-8	-	-	-	-	++	-	-	-	+++	++	++
Neu4Ac5Gc	α2-3	++	-	-	++	+++	-	-	-	++	-	-/+
	α2-3	+	-	-	+	+++	-	-	-	+++	-	-
Neu5Gc9Ac	α2-6	++	-	-	+++	+++	-	-	-	+	+	+
Neu5Gc ^{Me}		-	-	-	-	-	-	-	-	-	-	-
Ganglioside type		-	-	-	-	-	-	-	-	-	-	_°

726

727 Supplemental Figure S2: Summarized result of CD33 sialoglycan binding. The results of the 728 sialoglycan microarray binding of different wildtype and mutant CD33 proteins presented in Figure 3 are summarized here. A differential sialoglycan binding preference was observed when 729 730 wildtype and mutant human/chimpanzee CD33 proteins were tested on the microarray. Binding is 731 annotated with a positive (+) symbol and the strength of the binding is indicated by the number of 732 the symbols. +++ indicates a very strong binding. Negative (-) symbol implies non-binding and -733 /+ indicates very faint interaction. In some cases, only a few sulfated glycans showed strong 734 binding signal (indicated with asterisk). Degree (°) symbols indicate binding with a very few 735 numbers of glycans only. Linkage indicates the nature of the glycosidic bond of the terminal Sia 736 to the underlying glycan.

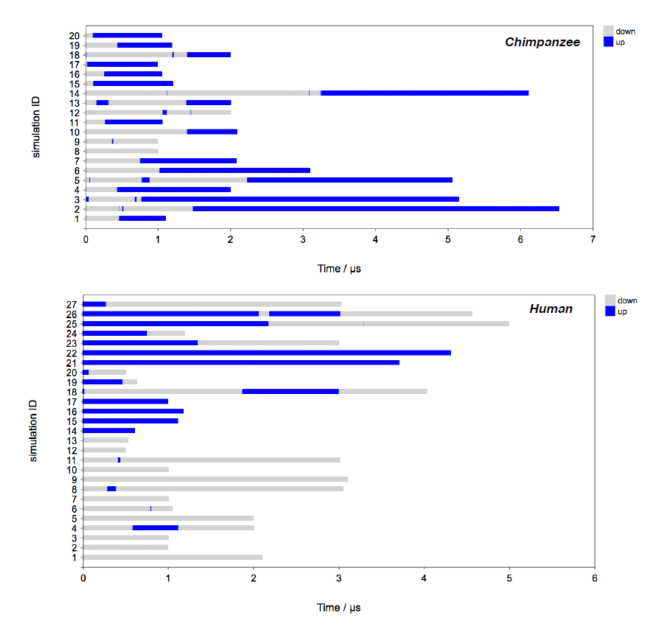
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737

- Supplemental Figure S3: Structure and dynamics of CD33. (A) Examples of x-ray structures of huCD33. PDB entries 5ibb (chain A: dark grey, chain B: white), 6d49 (lime). The dynamics of the C-C' loop and residues Phe21 and His45 are indicated. Positions of mutations present in chimpanzee are labeled on the pink spheres. (B) Amino acid sequences. 1: CD33 human (Uniprot),
- 742 2-3: PDB entries used for modeling. 3-4: sequences of the chCD33 models.

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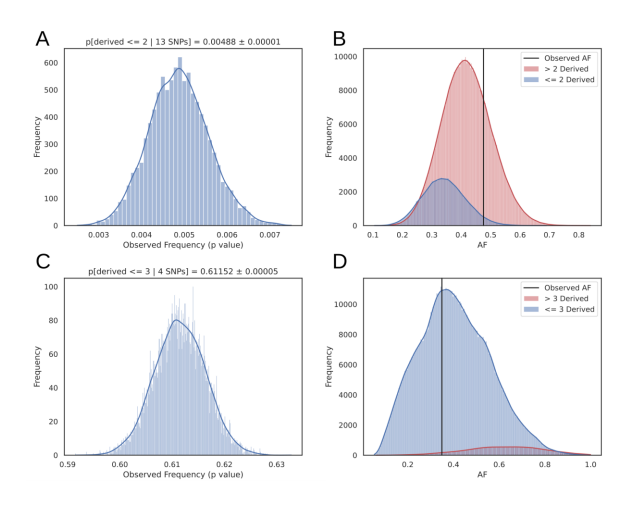


743

Supplemental Figure S4: MD trajectories of up/down states of Histidine at position 45 (His 45). Molecular dynamics of His45 side chain orientation. Individual MD trajectories of 'up'(blue)/'down'(grey) conformational states are shown. For chCD33, it can be observed reproducibly that simulations started with His45 in a 'down' conformational state undergo a transition to the 'up' conformational state on the microsecond timescale. Therefore, it may be concluded that chCD33 exists mainly with His45 in an 'up' orientation, which would be favorable for binding of Neu5Gc. For huCD33, both conformational states can exist for multiple µs, which

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- explains why huCD33 can bind to Neu5Ac (preferably binds when His45 is 'down') and Neu5Gc
- 752 (preferably binds when His45 is 'up').



753 754

755 Supplemental Figure S5: Resampling analysis of the 5.9 million SNPs from 1000 genomes 756 variants. As an alternative to matching AF directly, the set of filtered SNPs were further restricted 757 to those with a derived population frequency greater than 0.05 resulting in a universe of 5.9 million 758 SNPs. We estimated the probability of observing at most two SNPs derived in either the 759 Neanderthal or Denisovan reference genomes and a mean allele frequency as large as the empirical 760 variants of interest (AF = 0.476). By randomly drawing SNPs, we found that the probability of 761 observing 13 SNPs with such as high global allele frequency and lack of derived alleles in archaic 762 genomes to be highly unusual (*p*-value = 0.00487 ± 0.00001) (A). The low frequency is driven by 763 two factors, as shown in (B). Most of the SNPs sampled have more than two archaic-derived SNPs 764 (red curve). Of those with fewer than two archaic-derived SNPs, the overall allele frequency is 765 typically low compared to the target set. With other Siglec SNPs, resampling captures similar 766 properties (C and D), indicating the LOAD protective set does not represent a random sampling 767 from the genome.

APOELOAD, CVDrs7412, rs429358Encodes plasma protein APOE, is polymorphic in humans. Three alleles (E2, E3, E4) encode proteins with distinct affinity for lipoprotein particles. The ancestral E4 allele is associated with highest LOAD risk, and increased atherosclerosis and vascular dementia. The derived alleles E2 and E3 seems protective against LOAD, with the lowest risk is in homozygous E2 individuals.[4]PICALMADrs3851179 rs10792832Encodes phosphatidylinositol-binding clathrin assembly protein (PICALM), considered to be one of numerous reproducible risk genes for LOAD.[4]SPONIDementiars2618516Encodes the developmentally regulated protein F-spondin, reported to be a putative ligand for the amyloid precursor protein (APP).[6]TCFLC2Diabetesrs7903146Associated with impaired insulin secretion and enhanced hepatic glucose production.[6]ARID5BADrs2588969Gene encodes a member of AT-rich interaction domain (ARID) family of DNA binding proteins. The encoded protein forms a histone H3K9Me2 demethylase complex with PHD finger protein 2 and regulates the transcription of target genes involved in adipogenesis and liver development.[6]PILRAADrs1859788A cell surface inhibitory receptor that recognizes specific O-glycosylated proteins and expressed on various innate[6]	6, 47] [8, 65, [76] [49] [50]
APOELOAD, CVDrs7412, rs429358Three alleles (E2, E3, E4) encode proteins with distinct affinity for lipoprotein particles. The ancestral E4 allele is associated with highest LOAD risk, and increased atherosclerosis and vascular dementia. The derived alleles E2 and E3 seems protective against LOAD, with the lowest risk is in homozygous E2 individuals.[4]PICALMADrs3851179 rs10792832Encodes phosphatidylinositol-binding clathrin assembly protein (PICALM), considered to be one of numerous reproducible risk genes for LOAD.[4]PICALMADrs2618516Encodes phosphatidylinositol-binding clathrin assembly protein (PICALM), considered to be one of numerous reproducible risk genes for LOAD.[6]SPON1Dementiars2618516Encodes the developmentally regulated protein F-spondin, reported to be a putative ligand for the amyloid precursor 	76] [49]
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PILRA AD rs1859788 O-glycosylated proteins and expressed on various innate [,	[51]
immune cell types including microglia	[53]
CYP3A5Salt retention and hypertensionrs776746Cytochrome P450 (CYP) genes are abundant in animal, plant, and bacterial genomes and have evolved to metabolize a variety of diverse compounds.[.	[54]
PPARG Diabetes rs1801282 A nuclear hormone receptor that regulates adipogenesis []	[55]
BIN1ADrs7561528Also known as amphiphysin 2, has recently been identified as the most important LOAD risk locus[1]	[20]
SCG2 Hypertension rs1017448 Secretogranin II (SCG2) associates with hypertension [.]	[56]
CAPN10Diabetesrs2975760CAPN10 encodes a member of the calpain-like cysteine protease family that regulates blood glucose levels.[.	[57]
AGTSodium retentionrs699Sodium homeostasis links with hypertension[.	

LOAD: Late onset Alzheimer's disease; AD: Alzheimer's disease; CVD: Cardiovascular disease
 Supplemental Table S1: Genes affecting cognitive functions in post-reproductive age exhibiting
 disease-protective alleles uniquely in humans. The corresponding references for each of the genes

are mentioned in the table.

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Gene	SNP ID	Allele	Global frequency	African	East Asian	European	South Asian	American
CD33	rs12459419	С	0.789	0.949	0.814	0.69	0.84	0.52
		Т	0.211	0.051	0.186	0.31	0.16	0.48
	rs3865444	С	0.789	0.949	0.814	0.69	0.84	0.52
		Α	0.211	0.051	0.186	0.31	0.16	0.48
APOE	rs7412	С	0.925	0.897	0.9	0.937	0.96	0.96
		Т	0.075	0.103	0.1	0.063	0.04	0.04
	rs429358	Т	0.849	0.732	0.914	0.845	0.91	0.9
		С	0.151	0.268	0.086	0.155	0.09	0.1
		Т	0.351	0.105	0.407	0.371	0.39	0.39
PICALM	rs3851179	С	0.685	0.895	0.593	0.629	0.61	0.61
	rs10792832	Α	0.313	0.094	0.409	0.372	0.4	0.39
		G	0.685	0.895	0.593	0.628	0.6	0.61
SDOM	rs2618516	Т	0.341	0.259	0.302	0.382	0.52	0.24
SPON1		С	0.659	0.741	0.698	0.618	0.48	0.76
TCELCO	rs7903146	С	0.772	0.74	0.977	0.683	0.7	0.77
TCFLC2		Т	0.228	0.26	0.0023	0.317	0.3	0.23
ARID5B	rs2588969	С	0.532	0.472	0.482	0.641	0.63	0.42
		Α	0.468	0.528	0.518	0.359	0.37	0.58
PILRA	rs1859788	Α	0.341	0.102	0.612	0.321	0.29	0.5
		G	0.659	0.898	0.388	0.679	0.71	0.5
СҮРЗА5	rs776746	Т	0.379	0.82	0.287	0.05	0.33	0.2
		С	0.621	0.18	0.713	0.95	0.67	0.8
PPARG	rs1801282	С	0.93	0.995	0.974	0.88	0.88	0.88
		G	0.07	0.005	0.026	0.12	0.12	0.12
BIN1	rs7561528	G	0.8	0.809	0.881	0.683	0.87	0.74
		Α	0.2	0.191	0.119	0.317	0.13	0.26
SGC2	rs1017448	С	0.879	0.635	0.963	0.979	0.97	0.95
		Т	0.121	0.365	0.037	0.021	0.03	0.05
CADNIO	rs2975760	Т	0.882	0.971	0.907	0.841	0.79	0.87
CAPN10		С	0.118	0.029	0.093	0.159	0.21	0.13
AGT	rs699	Α	0.295	0.097	0.147	0.588	0.36	0.36
AGI		G	0.705	0.903	0.853	0.412	0.64	0.64

773

774 Supplemental Table S2: Analysis of Gene variants directly or indirectly affecting cognitive
775 function with their human population frequency. The global frequency of the SNPs identified in
776 Supplemental Table S1 was studied across different populations as indicated in the top of the
777 columns.

Amino	Human	Chimpanzee	Mutagenesis Primer
acid	CD33(M)_EK_Fc	CD33(M)_EK_Fc	Pairs Forward/Reverse $5' > 3'$
	. ,		rans_rorward/Reverse_3 > 3
position	Variant	Variant	
20	N20K	-	TGGATCCAAAaTTCTGGCTGCAAGTGCAGG
			TAGCCAGGGCCCCTGCCC
21	F21I	-	GGATCCAAATaTCTGGCTGCAAGTGCAG
			ATAGCCAGGGCCCCTGCC
22	W22R	-	TCCAAATTTCcGGCTGCAAGTGCAGG
			TCCATAGCCAGGGCCCCT
65	A65P	-	CCGGGAAGGAcCCATTATATC
			AACCAGTAACCATGAACTG
66	I66F	-	GGAAGGAGCCtTTATATCCAGG
			CGGAACCAGTAACCATGAAC
67	I67V	-	AGGAGCCATTgTATCCAGGGAC
			TCCCGGAACCAGTAACCA
69	R69G	-	CATTATATCCgGGGACTCTCCAGTG
			GCTCCTTCCCGGAACCAG
78	L78P	-	ACAAACAAGCcAGATCAAGAAGTACAGGAG
			GGCCACTGGAGAGTCCCT
96	P96L	-	CTTGGGGATCtCAGTAGGAACAAC
			GAGGCGGAATCTGCCCTG
148	L148V	-	GCCCAAAATCgTCATCCCTGG
			CTGTGGGTCAAGTCTGTC
152	T152A	-	CATCCCTGGCgCTCTAGAACC
			AGGATTTTGGGCCTGTGG
154	E154D	-	GCACTCTAGAtCCCGGCCACT
			CAGGGATGAGGATTTTGGG
21	-	I21F	GGATCCAAAAtTCCGGCTGCAAGTG
			ATAGCCAGGGCCCCTGTG
20	-	K20N	TGGATCCAAAtATCCGGCTGCAAGTGC
-			TAGCCAGGGCCCCTGTGG
L	1	1	

778

779 Supplemental Table S3: List of the mutagenesis primers used in the study to generate the CD33

780 mutants. Lowercase letters correspond to base change.

781 Supplemental File S1: List of the glycans used for the sialoglycan microarray. The complete

- 782 list of the chemoenzymatically synthesized glycans used to determine the binding profile of
- 783 different CD33 proteins are presented. The binding intensity of the different proteins (indicated on
- the top of the columns) towards the corresponding glycan are shown in the heatmap (same heatmap
- as in Figure 3). The red indicates maximum, and blue indicates minimum binding. R =
- propylamine linker present in the underlying glycan structure. Gal = galactose, GalNAc = N-
- acetylgalactosamine, Glc = glucose, GlcNAc = N-acetyl glucosamine, Fuc =L-fucose. The linkage
- 788 between the monosaccharides is indicated as α or β with numbers.

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