

Evolutionary conservation of human ketodeoxynonulosonic acid production is independent of sialoglycan biosynthesis

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Human metabolic incorporation of nonhuman sialic acid (Sia) *N*-glycolylneuraminic acid into endogenous glycans generates inflammation via preexisting antibodies, which likely contributes to red meat-induced atherosclerosis acceleration. Exploring whether this mechanism affects atherosclerosis in end-stage renal disease (ESRD), we instead found serum accumulation of 2-keto-3-deoxy-*D*-glycero-*D*-galacto-2-nonulosonic acid (Kdn), a Sia prominently expressed in cold-blooded vertebrates. In patients with ESRD, levels of the Kdn precursor mannose also increased, but within a normal range. Mannose ingestion by healthy volunteers raised the levels of urinary mannose and Kdn. Kdn production pathways remained conserved in mammals but were diminished by an M42T substitution in a key biosynthetic enzyme, *N*-acetylneuraminase synthase. Remarkably, reversion to the ancestral methionine then occurred independently in 2 lineages, including humans. However, mammalian glycan databases contain no Kdn-glycans. We hypothesize that the potential toxicity of excess mannose in mammals is partly buffered by conversion to free Kdn. Thus, mammals probably conserve Kdn biosynthesis and modulate it in a lineage-specific manner, not for glycosylation, but to control physiological mannose intermediates and metabolites. However, human cells can be forced to express Kdn-glycans via genetic mutations enhancing Kdn utilization, or by transfection with fish enzymes producing cytidine monophosphate-Kdn (CMP-Kdn). Antibodies against Kdn-glycans occur in pooled human immunoglobulins. Pathological conditions that elevate Kdn levels could therefore result in antibody-mediated inflammatory pathologies.

Introduction

Sialic acids (Sias) are a diverse family of 9-carbon backbone monosaccharides found in glycosidic linkage to outer tips of the glycan forest that covers all vertebrate cells and most secreted macromolecules (1–5). Given their widespread expression and localization, Sias have diverse roles in biology, evolution, and disease (2, 6–10). The 2 most common mammalian Sias are *N*-acetylneuraminic acid (Neu5Ac) and *N*-glycolylneuraminic acid (Neu5Gc). Neu5Gc is hydroxylated from Neu5Ac by cytidine monophosphate-*N*-acetylneuraminic acid hydroxylase (CMAH). Humans cannot synthesize Neu5Gc because of inactivation of the *CMAH* gene that occurred about 2–3 million years ago (11). We previously reported that most humans have circulating polyclonal antibodies against glycans bearing Neu5Gc (12, 13) and that anti-Neu5Gc-glycan

antibody generation correlates with the introduction of Neu5Gc during weaning (14). When exogenous Neu5Gc of dietary origin gets metabolically incorporated into sites such as epithelium and endothelium (15), glycans bearing this Sia act as foreign “xeno-autoantigens,” triggering inflammatory responses termed “xenosialitis” (13, 16–19). Disease-escalating effects of xenosialitis have been demonstrated in cancer and atherosclerosis models using human-like *Cmah*-null mice (20, 21).

2-Keto-3-deoxy-*D*-glycero-*D*-galacto-nonulosonic acid (Kdn) is another vertebrate Sia that was first discovered in the cortical alveolar polysialoglycoprotein of rainbow trout eggs (22) and is abundant in cold-blooded vertebrates. Although glycosidically conjugated Kdn was not originally found in mammals (23), free Kdn was detected in porcine submaxillary glands (23), cow milk products, human urine (24), and porcine milk (25). Elevated levels of free Kdn (and indirect evidence suggesting very low levels of conjugated Kdn) have also been reported in human ovarian (26) and throat cancers (27), and free cytosolic Kdn-containing *N*-glycans accumulate in human prostate cancers (28). Monoclonal anti-Kdn antibodies produced from Kdn-immunized mice also detected very small amounts of Kdn in rat pancreas (29, 30). Another

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monoclonal antibody against poly (2-8-linked) Kdn indicated the presence of this structure on megalin in the kidney (31–33), on cells of certain types of cancer (33), as well as on ceruloplasmin (34). However, to our knowledge, conclusive evidence for the occurrence of Kdn-glycans in normal human tissues is still missing.

Free Kdn is synthesized in the cytosol (refs. 35, 36 and Figure 1) from mannose 6-phosphate (Man-6P) (a C-2 epimer of glucose 6-phosphate) — which is itself used for the biosynthesis of *N*-linked glycans found on most secretory proteins — as a core component of glycosylphosphatidylinositol (GPI) anchors and as a metabolic precursor of fucose (37). Importantly, unlike many other free monosaccharides, mannose is very efficiently taken up by different cell types using a transporter that is insensitive to glucose (38, 39). Although specific mannose transporters have not been identified yet, mannose uptake suggests that endogenous mannose production may sometimes be inadequate for cellular requirements and that salvaging extracellular free mannose can be important.

Oral mannose administration can directly contribute to *N*-glycan synthesis (40), however, Man-6P is also endogenously derived from fructose-6-phosphate (Fru-6P) through phosphomannose isomerase (PMI) (ref. 39 and Figure 1). Inherited deficiency of PMI results in congenital disorder of glycosylation, type Ib (CDG-Ib) (PMI-CDG) associated with protein loss in intestines, liver disease, hypoglycemia, and blood-clotting disorders (41), and these patients respond well to oral mannose therapy (42). On the other hand, excessive mannose is toxic for honeybees (43, 44) and can blind baby PMI-KO mice (45). In both instances, there is a low ratio of PMI relative to hexokinase. This imbalance produces a metabolic flux, in which mannose is rapidly phosphorylated, but Man-6P cannot be efficiently converted to glycolytic intermediates. Thus, adenosine triphosphate (ATP) is wasted to produce a molecule without metabolic value to the cell, an effect that has been termed the “honeybee syndrome” (44, 46).

Although oral mannose is a potential therapeutic nutraceutical to treat certain types of CDG type I disorders (47), mannose supplements are also used to suppress urinary tract infections by *Escherichia coli* that bind mannose-rich glycans (48–50). Furthermore, recent murine studies showed that mannose supplementation has an immunosuppressive effect by inducing Tregs (51) and ameliorating high-fat diet-induced obesity and glucose tolerance by changing the microbiome profile (52). Several human studies revealed the association between increased plasma mannose levels and the risk of diabetes (53, 54), cardiovascular disease (55), and colorectal cancer risk (56). A recent study also suggested that mannose administration impairs tumor growth and enhances chemotherapy (57). However, to date, no comprehensive study has been performed to understand potential mannose toxicity in humans or the possibility of the regulation of physiological mannose homeostasis via conversion to Kdn.

Here, we report that, although serum-free mannose levels were increased but still maintained in an almost-normal range in patients with end-stage renal disease (ESRD) receiving hemodialysis, their free serum Kdn levels were substantially elevated. Furthermore, we show that humans not only excrete excess mannose into urine, but also metabolize it into free Kdn and excrete the Sia into urine, presumably to modulate mannose levels.

We also assess the evolutionary conservation (and/or convergent evolution) of genes encoding Kdn metabolic enzymes

and hypothesize that the conservation of Kdn biosynthesis helps to detoxify excess Man-6P and Fru-6P. We also show that man-nose-fed mammalian cells produced free Kdn in their cytosol, with predicted increases in certain genetic mutants, and that humans can have antibodies against glycosidically linked Kdn glycoconjugates (hereafter called Kdn-glycans).

Results

Increased circulating free Kdn in patients with ESRD. Previous studies in human-like *Cmah*-null mice showed that orally ingested glycosidically conjugated Neu5Gc can be metabolically incorporated into tissues, most prominently into epithelial and endothelial cells, driving antibody- and complement-mediated inflammatory progression of carcinomas (20) and atherosclerosis (21). In contrast, ingested free Neu5Gc is excreted rapidly into the urine of both rodents (58, 59) and humans (15). On the other hand, sustained, high-level exposure of cultured human cells to free Neu5Gc can result in incorporation that is sufficient to generate human anti-Neu5Gc-glycan antibody-mediated and complement-mediated inflammation (60). Thus, we hypothesized that accumulation of free Neu5Gc from the diet contributes to accelerated atherosclerosis in patients with ESRD. To explore this hypothesis, we examined free Sia profiles in sera collected from anonymized hemodialysis patients just prior to their weekly dialysis and compared them with samples from healthy volunteers. We did not see free Neu5Gc peaks in either ESRD or control sera, but instead observed a significant accumulation of other DMB-reactive α -ketoacids including Kdn and Neu5Ac in the ESRD samples (Figure 2, A–D, and Supplemental Figure 1; supplemental material available online with this article; <https://doi.org/10.1172/JCI137681DS1>). We confirmed the identity of Kdn and Neu5Ac peaks by liquid chromatography–mass spectrometry (LC-MS) (Supplemental Figure 2) using commercially available standards (Supplemental Table 1). Interestingly, we also noted a slight increase in 3-deoxy-D-manno-oct-2-ulose (Kdo), a bacterial octulosonic acid and component of plant polysaccharides, and other unknown peaks only in ESRD samples (Supplemental Figure 2). Serum-free Kdn levels in patients with ESRD were significantly elevated compared with levels in healthy volunteers (2.91 [0.68] μ M vs. 0.50 [0.25] μ M, mean [SD], $P < 0.001$; Figure 2B), as were the levels of serum-free Neu5Ac (16.9 [2.01] μ M vs. 2.34 [0.54] μ M, $P < 0.001$; Figure 2C). Although the levels of mannose (the metabolic precursor of Kdn) in patients with ESRD were significantly higher than those in healthy volunteers (83.8 [15.1] μ M vs. 44.5 [19.1] μ M, $P < 0.001$; Figure 2D), the levels in most patients with ESRD (10 of 16 patients) were within the normal range for humans (40–80 μ M) (61).

Ingested mannose is partially converted to Kdn and excreted into urine in humans. We next asked whether feeding mannose to healthy humans would result in increased Kdn production. In prior studies involving healthy humans, feeding mannose at 0.10–0.25 g/kg body weight resulted in a peak concentration of mannose in the blood of approximately 0.5 mM, with only mild gastrointestinal symptoms in 1 individual. However, at oral doses exceeding 0.25 g mannose/kg body weight, approximately half of the subjects reported mild gastrointestinal distress. No other symptoms were reported (47). We therefore studied healthy humans during

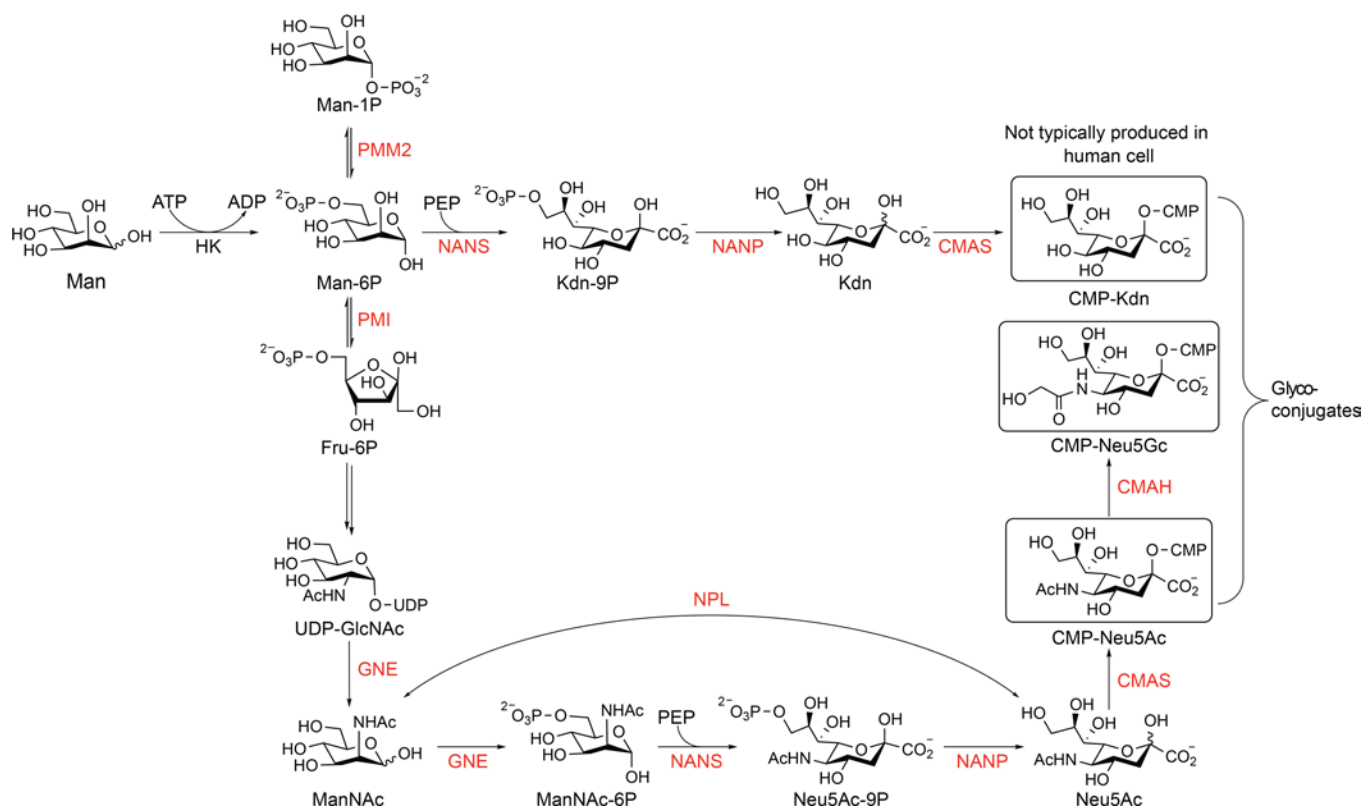


Figure 1. Metabolic pathway associated with mannose and Sias. Mannose is associated with Sia metabolism. Mannose is phosphorylated to Man-6P by hexokinase and ATP as a cofactor. Man-6P is converted to Kdn with NANS and *N*-acetylneuraminic acid phosphatase (NANP), and Kdn can be converted to CMP-Kdn with CMP *N*-acetylneuraminic acid synthetase (CMAS) in some species. Man-6P can also be transformed into Man-1P by phosphomannomutase 2 (PMM2). Man-6P is also converted to Fru-6P by PMI. Fru-6P is transformed into uridine diphosphate *N*-acetylglucosamine (UDP-GlcNAc), which is then converted to ManNAc-6P via GNE. ManNAc-6P is converted to Neu5Ac via NANS and NANP. NPL regulates Neu5Ac concentrations by mediating the reversible conversion to ManNAc and pyruvate, but human NPL does not work on Kdn. Neu5Ac is converted to CMP-Neu5Ac with CMAS, and CMP-Neu5Ac is converted to CMP-*N*-glycolylneuraminic acid (Neu5Gc) by CMAH. Humans cannot synthesize Neu5Gc because of inactivation of the *CMAH* gene about 2–3 million years ago.

mannose ingestion at 0.20 g/kg body weight. No obvious symptoms were reported, and serum mannose levels, measured by gas chromatography–mass spectrometry (GC-MS), rose 1 hour after ingestion to a maximum of 285 μ M (mean) and decreased subsequently within 8 hours of ingestion to 120 μ M (Figure 3A), as previously described (47). Urine mannose levels increased from 4 μ M (mean) (before mannose ingestion) to 11 μ M and then decreased to approximately 5 μ M (Figure 3A). Similarly, we observed a peak of free Kdn 4 hours after ingestion that remained at this higher level (Figure 3B). Overall, following the serum mannose peak at 1 hour, both mannose and free Kdn levels in urine rose within 4 hours of ingestion (Figure 3), suggesting that excess free mannose could be directly and indirectly (metabolized to free Kdn) eliminated by urinary secretion. Serum Neu5Ac and Kdn levels did not change after oral mannose administration, consistent with a previous animal study (36) and our findings of serum Kdn elevation only in patients with ESRD and not in healthy control individuals (Figure 2). To determine whether oral mannose administration results in Man-6P accumulation in human serum and urine, we measured Man-6P concentrations after 1 hour in the serum and after 4 hours in the urine of healthy humans. We could not detect any Man-6P using high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD), even after

mannose ingestion (Supplemental Figure 3). One possible explanation could be that Man-6P is not exported out of the cells. Another possibility is the presence of plasma/serum phosphatases that destroy Man-6P and that those enzymes do not exist in the urine. The latter possibility is supported by the observation that exogenously added Man-6P could only be detected in urine but not in serum samples (Supplemental Figure 3, C and F). Overall, these results suggest that mannose levels may be controlled directly by the urinary system and/or indirectly by Kdn metabolic pathways.

Dose-dependent mannose toxicity in cultured human and mouse cells. To our knowledge, there is no report confirming mannose toxicity in humans. Free Kdn is synthesized in the cytosol of mammalian cells fed 20 mM mannose (35, 36). However, in vitro and in vivo studies suggest that mannose toxicity in PMI-KO cells or embryos can be caused by Man-6P accumulation, which inhibits glucose metabolism and depletes intracellular ATP (39, 40, 46). To test mannose toxicity in human cells, we performed a continuous cell viability assay for mannose-fed cultures at varying concentrations (0–200 mM), using another monosaccharide, *N*-acetylglucosamine (GlcNAc) (which is not efficiently taken up), as a control for any hyperosmolar effect (Figure 4). Human embryonic kidney 293A (HEK293A) cells and human umbilical vein endothelial cells (HUVECs) were tested as representative cells of kidney epithelium and endothelium

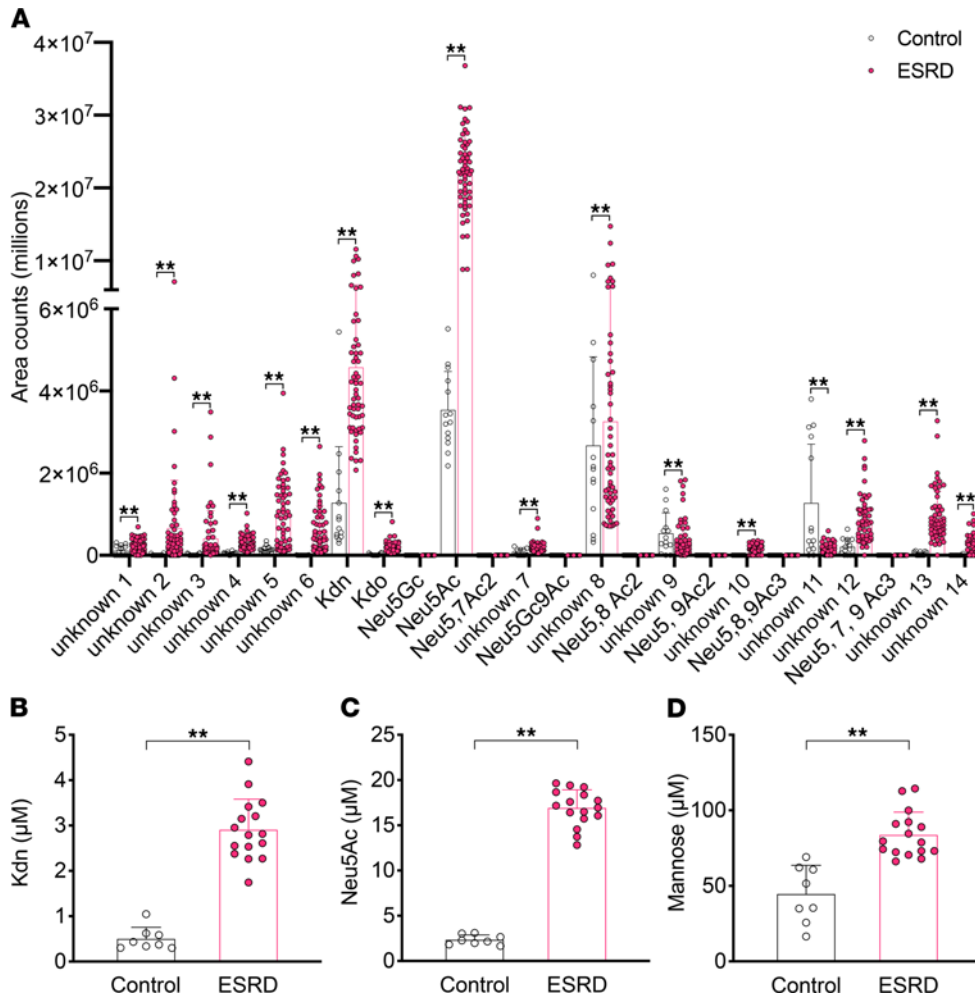


Figure 2. DMB-HPLC profiling of serum samples from patients on hemodialysis compared with serum from healthy individuals. Serum samples from (A) healthy volunteers (Control) ($n = 14$) and patients with ESRD on hemodialysis (ESRD) ($n = 58$) were analyzed by DMB-HPLC. Each dot represents each serum sample. Levels of the serum free (and CMP-) forms of (B) Kdn and (C) Neu5Ac were calculated with DMB-HPLC. (D) Serum mannose levels (normal range, 40–80 μM) were measured by GC-MS. Control ($n = 8$) versus ESRD ($n = 16$). Data are shown as the mean (SD). $**P < 0.01$, by 1-way ANOVA (A) and unpaired t test (B–D).

(25 μM), since this minimal mannose amount is indispensable for the viability of the PMI-KO cells.

Next, we sought to determine whether mannose feeding of the cultured cells could result in the production of cytosolic Kdn or whether the Sia upon formation is excreted into the growth media. To that end, we grew different human and mouse cells including HEK293A cells, HUVECs, BJAB K88 cells (GNE WT), and K20 cells (GNE-KO) as well as PMI WT and PMI-KO mouse embryonic fibroblasts in the presence of exogenous free mannose. On the basis of previous

encountering mannose in the blood stream, respectively. We also studied a human Burkitt's lymphoma cell line (BJAB K88, *GNE*^{+/+}, UDP-GlcNAc 2-epimerase/ManNAc kinase WT [GNE WT]) and a subclone deficient in Sia production (BJAB K20, *GNE*^{-/-}, GNE-KO) (62), as well as *PMI*^{+/+} (PMI WT) and *PMI*^{-/-} (PMI-KO) mouse embryonic fibroblasts (40). Neither HEK293A cells (Figure 4, A and B) nor HUVECs (Figure 4, C and D) showed differences between mannose and GlcNAc feeding, even at a high concentration (100 mM). However, the other cell lines clearly showed selective mannose toxicity at a lower concentration of 25 mM (Figure 4, E, G, and I) in comparison with GlcNAc feeding (Figure 4, F, H, and J). In particular, PMI-KO cells, when cultured at mannose concentrations higher than 1 mM (Figure 4K), displayed the “honeybee syndrome,” caused by overproduction of Man-6P, as previously shown (39, 40). A previous study confirmed that PMI-KO cells had Man-6P accumulation in the physiological condition (20–100 μM mannose) and that ATP depletion started when maximum levels of Man-6P had been reached (4–8 hours of culturing with 500 μM mannose supplementation) (40). We also observed a reduction in the viability of PMI-KO cells fed GlcNAc (10–30 mM) (Figure 4L) that was caused by a yet-unknown mechanism (63) and that could have been partly associated with ATP depletion during the GlcNAc feedings. Notably, unlike all other cells tested for the effect of GlcNAc, the PMI-KO cells in Figure 4L were always maintained in media supplemented with mannose

Kdn studies involving 20 mM mannose supplementation (35, 36) and our viability assay (Figure 4), we first tested 0, 1, 3, 5, 10, and 15 mM mannose supplementation and analyzed the concentration of Kdn relative to Neu5Ac in the cytosol (Supplemental Table 2) or in the growth media (Supplemental Table 3) by 1,2-diamino-4,5-methylenedioxybenzene-HPLC (DMB-HPLC). We decided to use 0 and 15 mM mannose supplementation conditions for all the human and PMI WT cells for further quantification ($n = 3$ experiments, Figure 5). Since PMI-KO cells failed to survive in the absence of mannose and required a minimal mannose concentration for their viability during the assay, we could not quantify Kdn in the mannose-depleted media for these cells. Instead, we used 25 μM (minimal concentration for cell survival), 250 μM (mimicking the maximum concentration in human sera, Figure 3A), and 1 mM mannose supplementation for Sia quantification in PMI-KO cells ($n = 3$ experiments, Figure 5). Analyses of purified free Sia and cytidine monophosphate (CMP-Sia) by DMB-HPLC confirmed that only CMP-Sia survived the reduction by NaBH₄ (ref. 64 and Supplemental Figure 4), allowing us to distinguish between free Sia and CMP-Sia in the cytosol. Most of the cytosolic Kdn represented the free form (NaBH₄ sensitive; up to 1000 pmole Kdn/million cells by DMB-HPLC) (Figure 5, A–E) and not CMP-Kdn (NaBH₄ resistant; less than 60 pmole Kdn /million cells) (Figure 5, F–J). Unlike the other human cell lines, which showed an increase in cytosolic accumulation of Kdn, cytosolic free (and

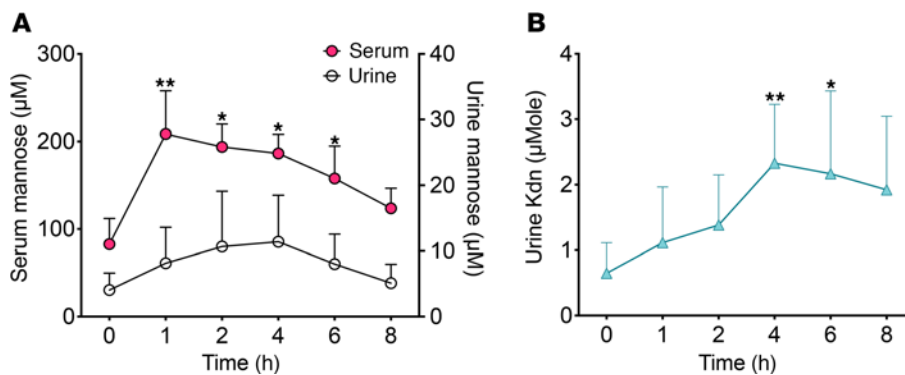


Figure 3. Human mannose ingestion test reveals increased urinary Kdn. Mannose (0.20 g/kg body weight) was used for the human ingestion test (healthy volunteers, $n = 5$). **(A)** Time course (0–8 hours after mannose ingestion) for serum and urine mannose levels, which were measured by GC-MS. **(B)** Urinary Kdn levels were evaluated by HPLC. Data are shown as the mean (SD). * $P < 0.05$ and ** $P < 0.01$, by 2-way ANOVA.

CMP-) Kdn levels in HEK293A cells did not change upon mannose feeding (Figure 5A). Instead, we found that Kdn levels in the media of HEK293A cells were elevated (Supplemental Table 3). We also analyzed an immortalized proximal tubule epithelial cell line from normal adult human kidney (HK2) (65), which showed a marked elevation of Kdn in the media after mannose feeding (Supplemental Table 3). Taken together with the Sia analyses in the cytosol and media, we conclude that human and mouse cells can produce free Kdn — and secrete it from cells — by an as-yet-unknown mechanism for the possible modulation of excess Man-6P.

Glycosidically conjugated Kdn rarely occurs in mammalian gly-cans. For a better understanding of the Kdn sialoglycome in eukaryotes, we searched available online databases for glycans with terminal glycosidically conjugated Kdn. Although glycosidically conjugated Neu5Ac and Neu5Gc were commonly observed in different eukaryotic groups, such Kdn-glycans were only reported in fish and amphibians, and not in any mammalian cells (Table 1). Indeed, in an extensive and in-depth mass spectrometric study of *N*-glycans from murine and human cell lines and tissues conducted by the Consortium for Functional Glycomics (66) (<http://www.functionalglycomics.org>), there was no evidence of Kdn-glycans in the approximately 425 human and mouse cell and tissue samples studied (Anne Dell and Stuart Haslam, Imperial College London, London, United Kingdom, personal communication). Taken together, these data suggest that the capacity for utilization of Kdn as a Sia conjugated to glycans was diminished or eliminated prior to the time that vertebrates became primarily land-based.

The Kdn biosynthetic pathway is conserved throughout vertebrate phylogeny and is under purifying selection in humans. As mentioned above, there is currently no conclusive evidence of Kdn-glycans in healthy mammalian tissues. On the basis of our observation of free Kdn in urine following mannose feeding (Figure 3B), we therefore asked whether human cells contain Kdn biosynthetic enzymes required for the production of Kdn from Man-6P. We investigated 7 human genes involved in mannose metabolism and conversion of Man-6P to Kdn (Figure 1) for evolutionary conservation, comparing them with their respective homologous genes in 5 species representatives of vertebrate phylogeny (Table 2).

While previous studies identified the conservation of Sia biosynthesis pathways in mammals (67), our results demonstrate that the enzymes capable of synthesizing Kdn from Man-6P are also conserved throughout vertebrate phylogeny and are experiencing ongoing purifying selection in humans and other vertebrate species (0.005–0.347, the ratio of the nonsynonymous substitution

rate [Ka] to the synonymous substitution rate [Ks], thus Ka/Ks, 64%–89% nt, and 54%–99% aa conservation) (Table 2). This finding further raises the question as to why Kdn biosynthesis is under purifying selection in organisms that do not appear to possess the capacity to utilize the monosaccharide in the synthesis of glycan structures. One explanation is that most of these genes are also involved in other conserved metabolic pathways. However, another nonmutually exclusive explanation for this selective pressure (i.e., preserving Kdn production) would be the necessity of these species to avoid the toxicity of excess Man-6P (see below).

A single aa in Sia 9-phosphate synthase that alters the rate of mannose to Kdn conversion is restored independently in 2 mammalian lineages, including humans. An earlier study showed that a single aa exchange (methionine 42 [M42] to any other aa except leucine) abolishes the ability of the human Sia 9-phosphate synthase (NANS) to synthesize Kdn-9P, while retaining its Neu5Ac-9P biosynthetic ability in vitro (68). We investigated the conservation of this key M42 position in other vertebrates. NANS protein sequences were highly conserved in a wide variety of lineages, e.g., mammalian species showed 87%–100% identity and 94%–100% homology, whereas nonmammalian vertebrates showed not less than 76% identity and 89% homology (Supplemental Table 4). We studied the phylogenetic topology including predicted ancestral aa and respective codons corresponding to M42 in human NANS (Figure 6 and Supplemental Figure 5). Considering the high homology to the human enzyme, we hypothesized that high Kdn-9P synthase activity was presumably associated with M42, also present in common ancestors. Remarkably, M42 was mainly found in non-mammalian vertebrates, e.g., aves, amphibia, and actinopterygii, in which, for the latter 2, the presence of glycosidically conjugated Kdn has been unequivocally demonstrated by MS analyses (Table 1). In mammals, M42 was found in monotremes (e.g., platypus) and metatheria (e.g., koala and opossum). In placentals, M42 was initially replaced by less active T42 but subsequently restored independently in 2 lineages: primates and artiodactyla (even-toed ungulates and cetaceans), apparently by convergent evolution. This finding again suggests an ongoing selection, possibly because of a need to modulate excess Man-6P into Kdn in clades that no longer use Kdn as a glycosidically conjugated Sia on glycoconjugates.

Mannose feeding in mutant cells can cause Kdn incorporation onto cell surfaces. Since healthy mammalian cells do not present glycosidically conjugated Kdn, we decided to include mutant cell lines in the analyses. On the basis of the cell viability assay results shown above, the highest concentration of mannose in cell

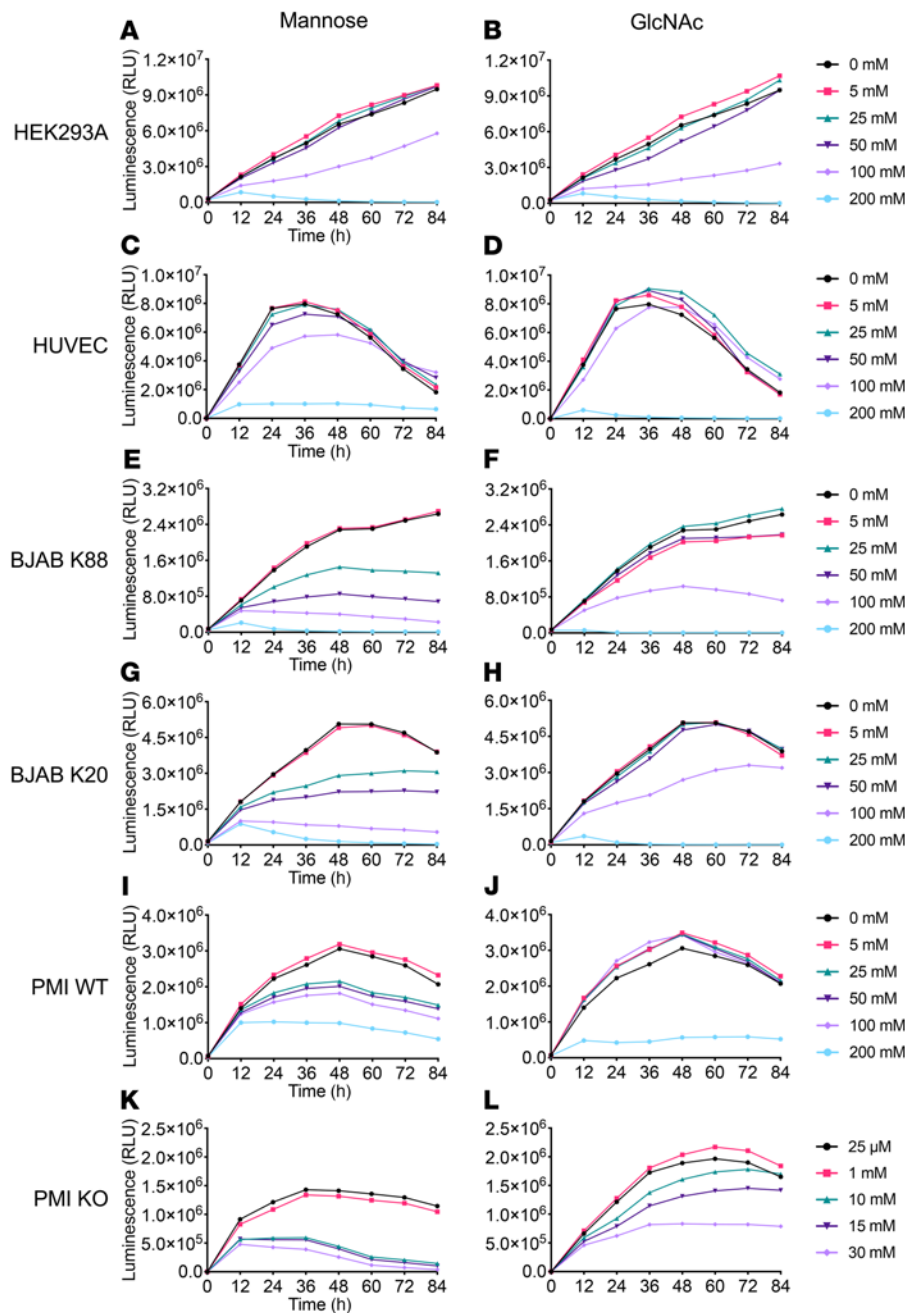


Figure 4. Toxicity of excess mannose in human and mouse cells in a dose-dependent manner. A continuous cell viability assay (0–84 hours) was performed to evaluate mannose toxicity to various human and mouse cell lines including mutants of Kdn-associated metabolic enzymes. The mannose feeding concentration was set at 0–200 mM, and the same concentration of GlcNAc was used as a control for the hyperosmolarity effect. Results shown are for (A and B) HEK293A cells; (C and D) HUVECs; (E and F) BJAB K88 cells (GNE WT); (G and H) BJAB K20 cells (GNE-KO); (I and J) PMI WT cells; and (K and L) PMI-KO cells. (L) PMI-KO cells were always maintained in mannose-supplemented (25 μ M) media for their survival during GlcNAc feeding. Data are representative of 2 independent experiments.

culture media was determined to be 15 mM (1 mM for PMI-KO cells) to maintain optimal cell viability. As mentioned earlier, the total amount of free (and CMP-) Kdn in cytosol or media was elevated in all types of cells after mannose supplementation (Figure 5 and Supplemental Tables 2–6), but no significant increase in the concentration of CMP-Kdn (Figure 5).

To evaluate the possible production of Kdn-containing glycan structures, we analyzed the Sia content of various cell lines fed with mannose and treated with NaOH and NaBH₄ to destroy any unconjugated Sia, and then examined by DMB-HPLC for glycosidically conjugated Sias from the total cell lysate (Supplemental Table 4). The highest Kdn/Neu5Ac ratio was observed in BJAB K20 cells, which, because of a GNE mutation, cannot make internal Neu5Ac. In line with this, BJAB K20 cells also showed a

high amount of free Kdn in the cytosol as well as in the media (Supplemental Tables 2 and 3). However, total conjugated Kdn levels were much lower than free and CMP-Kdn levels in the cytosol and media, even in BJAB K20 cells (Supplemental Tables 2–4). We also collected membrane fractions after ultracentrifugation, and DMB-HPLC analyses showed conjugated Kdn in the membrane fraction, especially in BJAB K20 and PMI-KO cells (Supplemental Table 5). Next, we used 2 previously reported anti-Kdn-glycan monoclonal antibodies, kdn3G (69) and kdn8kdn (70), which have specific affinities for glycans containing α 2-3- or α 2-8-linked Kdn, respectively, but not for α 2-6-Kdn-glycan structures (29, 69). Flow cytometry using these anti-Kdn antibodies did not detect any peak shift for BJAB K20 cells (Figure 7, A and B), probably because BJAB K20 cells mainly form α 2-6-linked Sia that are undetectable with the 2 antibodies used for the assay (71, 72). In contrast to BJAB K20 cells, PMI-KO cells presented small amounts of Kdn in α 2-3 and α 2-8 linkage on cell-surface structures upon feeding with mannose (Figure 7, C and D).

Fish Cmas gene expression plus mannose feeding can force Kdn incorporation onto some normal cell surfaces. Prior to transfer to glycans, Sias are activated into CMP-Sias, a reaction involving cytidine triphosphate (CTP) and catalyzed by CMP-Sia synthase, encoded by the CMP *N*-acetylneuraminic acid synthetase (CMAS) gene. As we showed earlier, Kdn incorporation into cell-surface glycans seems to be rare in comparison with free Kdn production, even after mannose feeding, probably because human CMAS and mouse CMAS preferentially work on Neu5Ac (73–75). Indeed, human CMAS has 94% identity to the murine enzyme (refs. 75, 76 and Supplemental Table 7), and the activity of the recombinant murine CMAS was 15 times lower for Kdn than for Neu5Ac (75). Our investigation into the

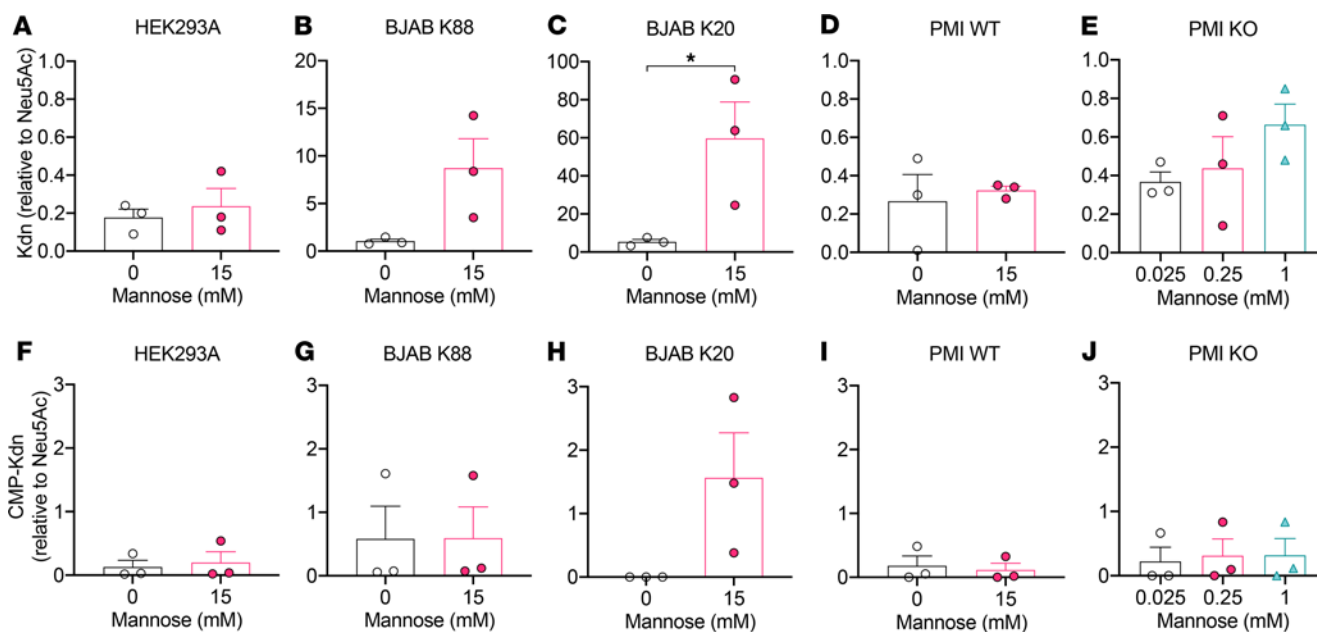


Figure 5. The Kdn/Neu5Ac ratio increases in cytosolic fractions after mannose feeding in human and mouse cell types. Levels of cytosolic free (and CMP-) Kdn (A–E) and CMP-Kdn (F–J) were measured by DMB-HPLC. Results are shown for (A and F) HEK293A cells; (B and G) BJAB K88 (GNE WT) cells; (C and H) BJAB K20 (GNE-KO) cells; (D and I) PMI WT mouse embryonic fibroblasts; and (E and J) PMI-KO cells. CMP-Sias were measured by DMB-HPLC after NaBH₄ treatment. The mannose concentration was set at 0 or 15 mM, except for PMI-KO cells (0.025, 0.25, and 1 mM). Data are shown as the mean (SD). **P* < 0.05, by unpaired *t* test.

phylogenetics of these pathways brought attention to a whole-genome duplication event that occurred in the teleost lineage of fish (77). This duplication produced 2 fish *Cmas* paralogs known as *Cmas1* and *Cmas2* (78). Since our previous results showed that HEK293A cells mainly make and excrete excess free Kdn into media, recombinant zebrafish *Cmas* plasmid vectors were transfected into HEK293A cells (Figure 7E, 30%–40% of cells were transfected, by checking transfection efficiency using an anti-Flag antibody, and data not shown). Flow cytometry using anti-Kdn antibodies showed no significant peak shift in *Cmas*-transfected HEK293A cells fed with up to 15 mM mannose (data not shown). However, DMB-HPLC analysis of the membrane fraction showed that zebrafish *Cmas* transfection could result in glycosidically conjugated Kdn (exemplarily shown for *Cmas2*-transfected HEK293A cells in Figure 7, H and I). The ratio of glycosidically conjugated Kdn to Neu5Ac, especially in mannose-fed samples, was lower in *Cmas1*-transfected cells (Supplemental Table 6), consistent with differential substrate specificity in vitro (78). Interestingly, the amount of conjugated Neu5Gc (traces derived from FCS) was lower than Neu5Ac but still higher than Kdn in human cells (Figure 7, F–I), confirming a limited utility for Kdn compared with Neu5Gc in glycosylation.

Humans have circulating polyclonal antibodies against Kdn-glycans. We have previously shown that humans have varying levels of circulating antibodies against glycans bearing the nonhuman Sia Neu5Gc (13, 14, 79). Although glycosidically conjugated Kdn is not reported in normal human tissues, some malignant cells may have the potential to incorporate Kdn on their surface sialoglycans in a Kdn-rich environment. To probe for the presence of any circulating anti-Kdn-glycan antibodies, we used chemoenzymatically synthesized biotinylated glycan containing Kdn α 2-3 linked to Gal β 1-4GlcNAc (Figure 8A) to screen commercially available pooled normal

human sera in an ELISA-based assay and found the presence of IgG antibodies against this Kdn-glycan epitope (Figure 8B). We also used Neu5Ac and Neu5Gc linked sialoglycans alongside non-sialylated glycans containing the same underlying Gal β 1-4GlcNAc epitopes to determine the presence of antibodies against those corresponding epitopes (Figure 8, A and B). While our current data do not present the complete repertoire of anti-Kdn-glycan antibodies in humans, they provide evidence for the selective occurrence of such antibodies in normal human sera.

Discussion

Mannose levels in mammalian sera are variable but seem to be controlled within a restricted range (20–160 μ M) (61). Levels of mannose can be affected by ingestion of mannose-rich food such as fruits, vegetables, yeasts, etc. (46, 80), by lysosomal release by *N*-glycan processing (61, 81), and by recovery and reabsorption by sodium glucose cotransporters (SGLTs) (82). SGLTs may work as mannose transporters in both intestine and kidney (82, 83). Mannose is also transported into mammalian cells via glucose transporters, but mannose-specific transporters have not yet been identified (84). Almost all transported mannose (95%–98%) is catabolized via PMI, and in [³H]-mannose radioactive metabolic labeling experiments, only up to 2% of mannose taken up is used for *N*-glycosylation (61, 81). Interestingly, a later study using more sensitive ¹⁵C-glucose and ²H-mannose isotopic labeling and GC-MS analysis showed that transported mannose is utilized more efficiently than glucose: while 1.8% of transported mannose appears in *N*-glycans, only 0.026% of transported glucose is utilized in *N*-glycan synthesis (85).

Furthermore, through a nocodazole-sensitive transporter, mannose from lysosomal *N*-glycan processing is transported out of the cell as free mannose without being catabolized (81). The

Table 1. Results of glycan database searches for glycosidically linked Kdn

Database	Total number of glycan structures	Occurrence of Neu5Ac	Glycans with Neu5Ac	Glycans with Kdn	Occurrence of Kdn	Ref.
GlyGen (https://www.glygen.org)	29,290	Mammals	8963	None	(99)	
GlyConnect (https://glyconnect.expasy.org)	3435	Mammals, fish, aves, amphibians	371	35	Fish, amphibians	(100)
UniCarb-DB (https://unicarb-db.expasy.org)	1118	Mammals	415	None	(101)	
LipidBank (http://lipidbank.jp)	7009	Mammals, fish, aves, amphibians	100	17	Fish, amphibians	(102)
GlyTouCan (https://glytoucan.org)	117,832	Mammals, fish, aves, amphibians	6570	334	Fish, amphibians	(103)

Table 2. Conservation of human Kdn biosynthesis pathway genes

	<i>PMM2</i>	<i>PMI</i>	<i>NPL</i>	<i>NANS</i>	<i>NANP</i>	<i>GNE</i>	<i>CMAS</i>
Ka/Ks							
<i>Homo</i>	0.000	0.000	0.000	0.000	0.000	0.000	0.163
<i>Mus</i>	0.047	0.132	0.100	0.029	0.069	0.005	0.317
<i>Gallus</i>	0.115	0.329	0.265	0.038	0.166	0.023	0.347
<i>Anolis</i>	0.162	0.276	0.274	0.089	0.292	0.068	0.201
<i>Xenopus</i>	0.206	0.265	0.324	0.151	0.272	0.091	
<i>Danio</i>	0.176	0.279	0.337	0.125	0.397	0.113	^A 0.038, ^B 0.264
% aa							
<i>Homo</i>	100	100	100	100	100	100	100
<i>Mus</i>	92	85	87	95	94	99	94
<i>Gallus</i>	85	64	71	87	72	93	77
<i>Anolis</i>	79	67	69	85	68	92	84
<i>Xenopus</i>	69	62	54	78	59	86	63
<i>Danio</i>	73	63	57	79	50	81	^A 60, ^B 53
% nt							
<i>Homo</i>	100	100	100	100	100	100	100
<i>Mus</i>	88	87	86	89	86	88	88
<i>Gallus</i>	78	70	74	81	67	81	76
<i>Anolis</i>	75	71	71	77	72	80	78
<i>Xenopus</i>	67	67	64	74	74	77	68
<i>Danio</i>	71	69	66	74		71	^A 68, ^B 75

Ka/Ks (top), percentage of aa identity (middle), and percentage of nt identity (bottom), values of genes involved in the synthesis of Sia across the vertebrate phylogeny: *Homo sapiens* (human); *Mus musculus* (mouse); *Gallus gallus* (chicken); *Anolis carolinensis* (green anole lizard); *Xenopus laevis* (African clawed frog); *Danio rerio* (zebrafish). For zebrafish: ^A*Cmas1* gene and ^B*Cmas2* gene. The whole-genome duplication event that affected *Cmas* in Teleostei also produced a paralog of *Nans*; however, the expression and function of the *Nansb* gene are unknown, so for this phylogenetic analysis, we focused on zebrafish *Nansa* (96).

reason for these separate pathways for intracellular mannose is unknown, but cells may need to eliminate excess mannose to avoid accumulation of toxic products such as Man-6P. Indeed, mannose toxicity was confirmed in our cell viability assay most prominently in PMI KO cells (Figure 4). High levels of mannose are also teratogenic in rat embryo cultures and pregnancy models, as a result of inhibition of glycolysis (86, 87).

Free Man-6P might be directly metabolized into Kdn or through the hexosamine biosynthetic pathway (Figure 1). Although human cells can synthesize Kdn from Man-6P, Kdn is not (or only in very small quantities) incorporated into glycoconjugates in healthy

humans. Given all these observations, we hypothesize that Kdn synthesis is conserved in mammalian metabolism to buffer cellular levels of Man-6P. Concordantly, our phylogenetic sequence analysis suggests an underlying evolutionary pressure to conserve the metabolic pathway that converts mannose to Kdn in vertebrates (Table 2). However, the capability to incorporate Kdn into glycoconjugates differs between vertebrate classes. The apparent lack of Kdn in the glycan determinants of mammals contrasts strikingly with the previously mentioned examples of Kdn-containing structures in the gametes and mucus secretions of amphibia and fish and suggests that the enzymes necessary for Sia incorporation may have lost their

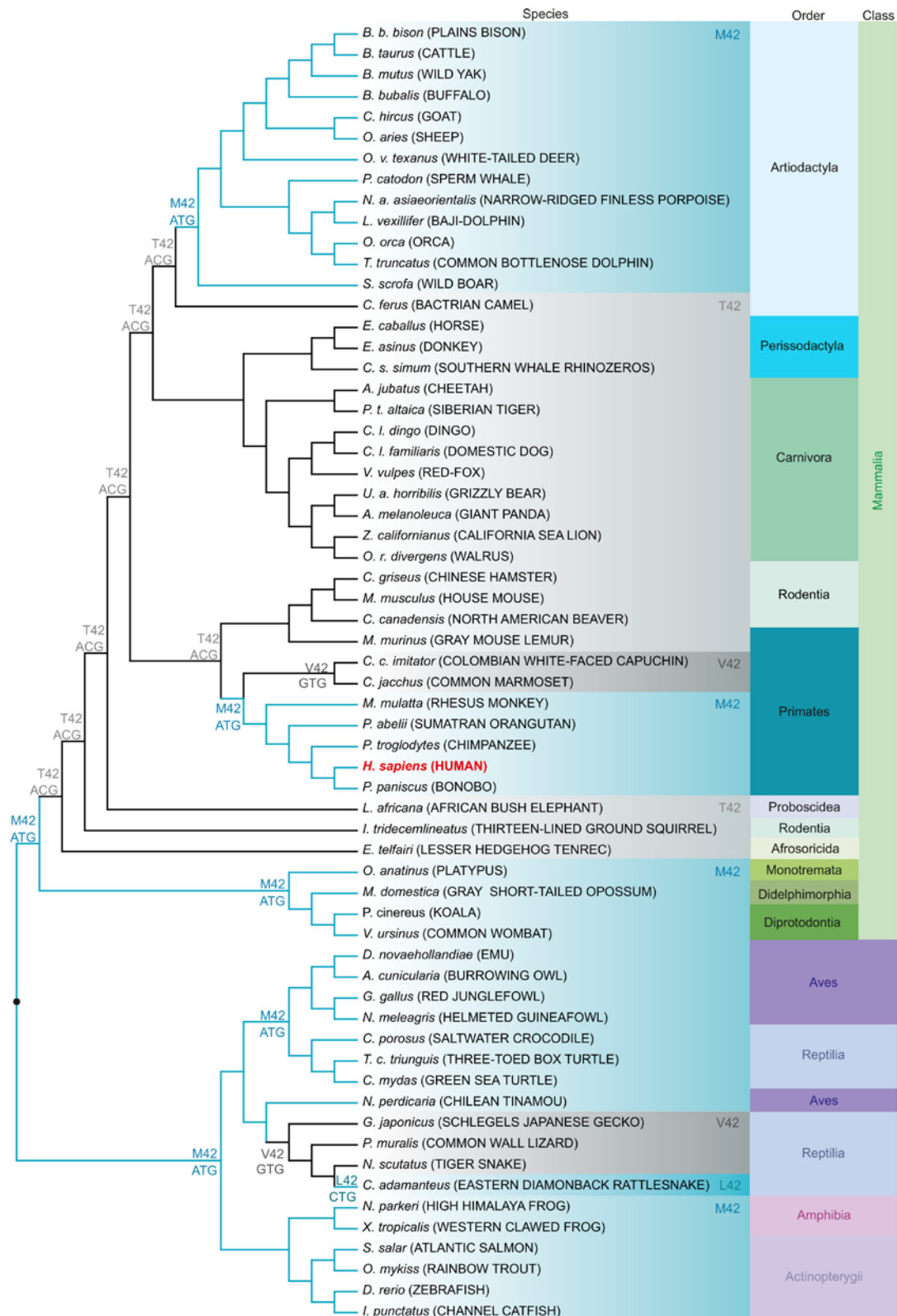


Figure 6. A single aa in Sia 9-phosphate synthase that alters the rate of mannose-to-Kdn conversion is restored independently by convergent evolution in 2 mammalian lineages including humans. Phylogenetic tree topology of vertebrate NANS nt sequences obtained using the maximum likelihood method. The tree was rooted on nonmammalian vertebrates (●) and is not drawn to scale. Human NANS with methionine or leucine at position 42 shows Kdn-9P synthase activity. Species with methionine or leucine at the position aligned to M42 in human NANS are shaded in light blue, and species with threonine (T42) or valine (V42) are shaded in light and dark gray, respectively. Predicted nt encoding the aa at the respective position in the ancestors are given at the nodes. Human NANS is shown in red. Latin species names are italicized and common names are given in uppercase letters. Mammals are further classified in orders and all vertebrates in classes.

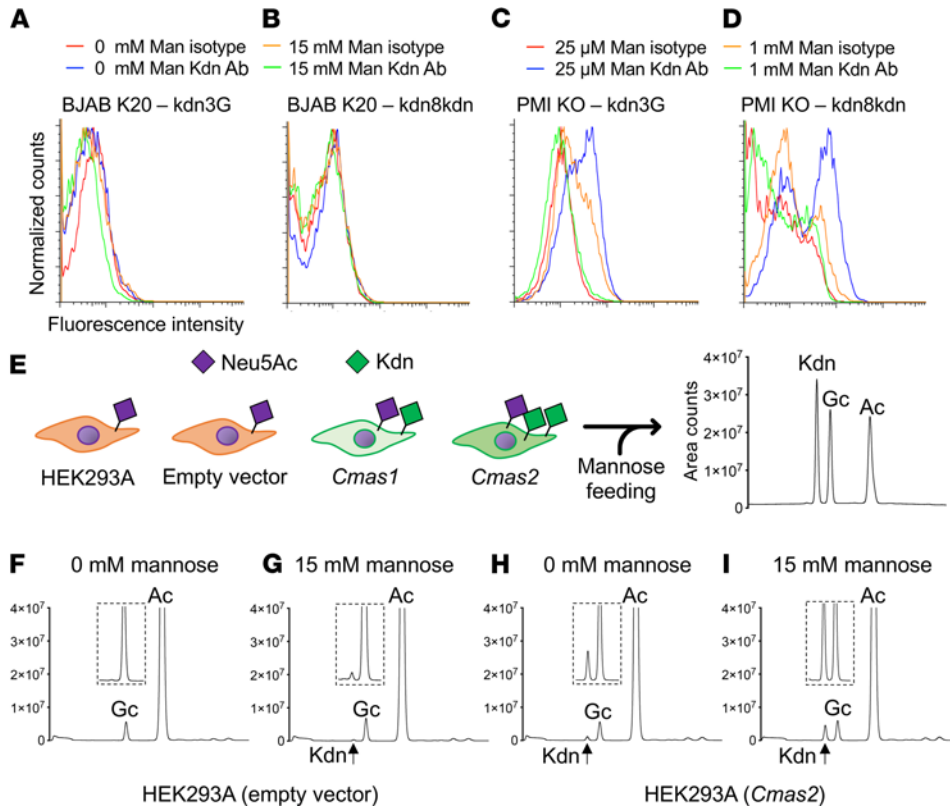


Figure 7. Gene mutations or fish *Cmas* transfection can enforce production of the conjugated form of Kdn. Mouse monoclonal anti-Kdn IgG antibody, with kdn3G recognizing the (Kdn) ganglioside GM3 and mouse monoclonal anti-Kdn IgM antibody, with kdn8kdn recognizing the α 2,8-linked Kdn were used for flow cytometry to analyze cell-surface expression of Kdn. Results are shown for (A and B) BJAB K20 (GNE-KO) cells and (C and D) PMI-KO cells, with or without mannose feeding. (E) Empty vector, *Cmas1*, and *Cmas2* were transiently expressed in HEK293A cells. Cells were harvested after 36 hours of mannose feeding (0 or 15 mM), and then membrane fractions were prepared to analyze the conjugated form of Sia by DMB-HPLC analysis (Kdn, Neu5Gc, and Neu5Ac). Representative HPLC peaks for empty vector (F and G) or *Cmas2*-transfected (H and I) membrane samples. (F and H) 0 mM and (G and I) 15 mM mannose feeding. Data are representative of 2 independent experiments. Ac, Neu5Ac; Gc, Neu5Gc; Man, mannose.

activity toward Kdn since the relatively ancient divergence of warm- and cold-blooded vertebrates.

For a free monosaccharide to be incorporated into a glycan structure, it must first be activated to a high-energy donor, e.g., conjugated with a nt, and then transferred to a glycosyl acceptor. In humans, CMAS and sialyltransferases (STs) retain the enzyme promiscuity necessary to efficiently conjugate exogenous Neu5Gc to CMP-Neu5Gc and subsequently incorporate the xenogenic Sia into a “xeno-autoantigen” glycan structure. However, the pseudogenization of the human *CMAH* gene occurred relatively recently, after the divergence of humans and chimpanzees. Compared with Neu5Gc, the efficiency of Sia in activating and transferring enzymes toward Kdn seemed to be much lower, since we only detected trace amounts of glycan-conjugated Kdn in human and mouse cells cultured in high-mannose media compared with the efficient incorporation of Neu5Gc present in trace amounts in the FCS-supplemented media. In contrast, Kdn is found in abundance in fish glycoconjugates based on a *Cmas* paralog that has evolved preferential activity toward Kdn. Using a previously published expression construct (78), we show here that in vitro expression of the Kdn-active zebrafish *Cmas* enzyme forced the expression of small amounts of Kdn-containing cell-surface glycans also in HEK293A cells (Figure 7 and Supplemental Table 6), which normally excreted excess Kdn to maintain a constant cytosolic Kdn concentration (Figure 5 and Supplemental Table 3).

Free Neu5Ac or Kdn is synthesized by NANS from *N*-acetylmannosamine (ManNAc) or Man-6P and phosphoenolpyruvate, respectively. In vitro analyses showed that the biosynthetic activity of Kdn is based on methionine or leucine at position 42 in human NANS. In line with glycosidically conjugated Kdn, amphibia and

fish NANS harbors methionine at position 42 (M42). Mouse NANS harbors the less active threonine (T42) and shows low Kdn biosynthesis in vitro (74). However, synthesis of free Kdn was observed in mouse cell culture in a high-mannose media, indicating residual enzymatic activity on Man-6P. Phylogenetic analyses showed that the common vertebrate ancestor harbors the Kdn-metabolizing M42, which is maintained mainly in nonmammalian classes and probably got lost during placentation in mammals (Figure 6). Reversion to methionine in placentals occurred independently by convergent evolution in primates and artiodactyla. Given these phylogenetic occurrences of Kdn biosynthetic enzymes, it is tempting to speculate that these orders might be more prone to a mannose-rich diet and/or lack microbiota capable of mannose uptake. Free Sias are catabolized by action of *N*-acetylneuraminidase pyruvate lyase (NPL) in vivo. However, human NPL has a 5-fold lower activity on Neu5Gc compared with Neu5Ac and no detectable activity on Kdn (88), thus favoring the persistence of free Kdn.

Interestingly, the distribution of Kdn in tissues of fish such as Atlantic salmon vary between skin and intestinal mucins; *O*-glycan structures showed that Neu5Ac, Neu5Gc, and Kdn are distributed in skin mucin, however, intestinal mucin contains only Neu5Ac (89, 90). Perhaps different host-pathogen interactions between aquatic and terrestrial habitats and/or constant exposure to hydrophilic aqueous environments produced different evolutionary pressures on Sia presentation.

In this study, we explored why the conversion of mannose into Kdn has been conserved in mammals like humans, despite the lack of conclusive evidence of Kdn presence in sialoglycoconjugates. On the basis of our human experiments, we hypothesize that mannose is handled like glucose in the kidney, being reabsorbed in the tubules,

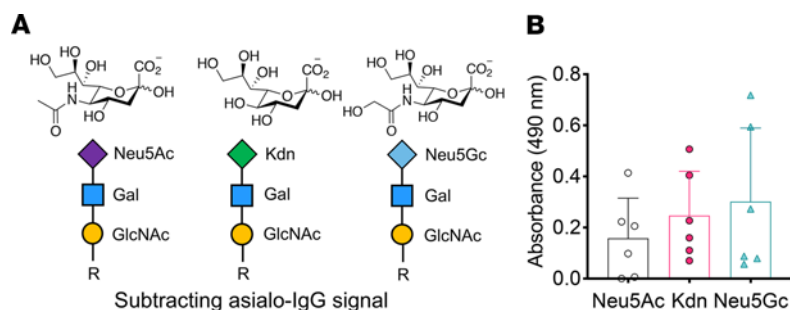


Figure 8. ELISA of anti-sialoglycan antibodies in human sera. (A) The chemoenzymatically synthesized, defined biotinylated glycans Neu5Ac α 2-3Gal β 1-4GlcNAc-, Neu5Gc α 2-3Gal β 1-4GlcNAc-, Kdn α 2-3Gal β 1-4GlcNAc-biotin, respectively. (B) Anti-Sia antibodies (Neu5Ac, Kdn, and Neu5Gc) were detected in pooled normal human sera (IVlg) samples ($n = 6$ each). The absorbance of the wells coated with Gal β 1-4GlcNAc- (negative control) glycan was subtracted from the corresponding sialoglycan-containing wells to determine the relative absorbance resulting from the specific anti-Sia antibodies. Data are shown as the mean (SD), and statistical significance was determined by 1-way ANOVA.

as previously reported (83, 91, 92), which implies that mannose is biologically “valuable.” However, excess mannose could also be excreted into urine, just as occurs with excess glucose in patients with diabetes (direct mannose control, Figure 3A). Moreover, phylogenetic analysis of vertebrate Kdn synthesis, together with *in vivo* oral mannose administration and *in vitro* mannose feeding studies, suggested that excess exogenous mannose is transported into the cytosol, then metabolized to Kdn through the conserved metabolic pathway, excreted from the cells, and finally eliminated from the body through the urine (indirect mannose control). We confirmed by mannose supplementation in the media of different human (HEK293A, HUVEC, BJAB, HK2) as well as mouse (PMI) cell lines that mammalian cells very efficiently excreted cytosolic Kdn into the media (Supplemental Table 3). Furthermore, Man-6P was not detectable in human serum or urine even after mannose ingestion (Supplemental Figure 3), consistent with our hypothesis that Kdn synthesis helps minimize the buildup of toxic Man-6P in the cytosol under conditions of mannose excess. However, we speculate that this is not the only reason. For example, mannose is 5 times more reactive in detrimental nonenzymatic protein glycation than is glucose (93) and therefore needs to be regulated. Also, we recently found that free Kdn can compete with host Neu5Ac for uptake by commensal and/or pathogenic bacteria, affecting their interaction with host immunity (94).

Overall, our work can help explain the evolutionary conservation of Kdn biosynthesis throughout the higher vertebrate phylogeny. Even though higher vertebrates, including humans, do not utilize Kdn in glycoconjugate production, Kdn synthesis remains a critical metabolic pathway that buffers the potentially toxic concentration of cytosolic Man-6P and Fru-6P. Moreover, humans produce anti-Kdn antibodies, probably because of the presentation of Kdn-containing antigens originating from symbiotic flora, as has been observed with human anti-Neu5Gc antibodies (14, 94). Our findings regarding the Kdn/mannose pathway (Figure 1) may contribute to identifying the exact pathways by which Kdn is taken up into cells and tissues and by which humans produce anti-Kdn antibodies. Ultimately, deciphering the role of Kdn metabolism may lead to a better understanding of human diseases, including

congenital disorders of glycosylation involving PMI, NANS, and NPL (88, 95, 96), as well as ESRD. It also remains possible that small amounts of conjugated Kdn may appear in tissues under pathological conditions like ESRD with high free Kdn levels and mediate inflammatory reactions with circulating anti-Kdn-glycan antibodies. If human cells are forced to incorporate small amounts of Kdn into surface glycans (e.g., under conditions of excess Kdn accumulation such as in acute or chronic uncontrolled renal failure), low levels of cell-surface accumulation could result in reactivity with such antibodies.

Methods

Human studies involving serum samples from patients with ESRD and healthy volunteer controls. Sera from patients with ESRD were collected just prior to weekly dialysis. Control samples were collected from healthy volunteers. All samples were anonymized, numbered, and kept at -80°C before analysis.

Human mannose ingestion test. Five informed and consented healthy volunteers ingested 0.2 g mannose/kg body weight dissolved in 500 mL drinking water, as previously tested (47). Blood samples (up to total 60 mL/person) were collected by venipuncture at 0, 1, 2, 4, 6, and 8 hours. Urine samples were collected as near as possible to the aforementioned times. All samples were analyzed for free and conjugated Kdn, Neu5Gc and Neu5Ac, as well as for mannose.

Cell culture and transfection. HEK293A cells (Thermo Fisher Scientific) were cultivated in RPMI 1640 media with 10% FBS; HUVECs (American Type Culture Collection [ATCC]) in EBM2 media (Lonza); BJAB K20 and BJAB K88 cells (provided by Stephan Hinderlich and the late Werner Reutter, Beuth University of Applied Sciences, Berlin, Germany) in RPMI 1640 media with 10% FBS; PMI WT and PMI-KO cells in DMEM containing 1 g/L glucose and 10% FBS; and HK2 cells in keratinocyte serum-free media supplemented with bovine pituitary extract and recombinant EGF (all from Gibco, Thermo Fisher Scientific). Transient transfection of *Cmas1* and *Cmas2* (and empty vector) was performed using polyethylenimine (PEI) at a final concentration of 18 $\mu\text{g}/100$ cm tissue culture dish and 8 μg plasmid DNA following standard protocols. The transfection media were removed after 4.5 hours, and the cells were incubated in complete media with or without added mannose at the indicated concentrations.

Cell viability measurement. Cell viability was measured using the RealTime-Glo MT Viability Assay (Promega) according to the instructions manual. Cells were incubated in culture media at various concentrations (25 μM –200 mM) of mannose or *N*-acetylglucosamine, which was used as a control for osmolarity effects. MT Cell Viability Substrate and NanoLuc enzymes (both from Promega) were also added to the culture, and viability was monitored by reading luminescence at 0, 12, 24, 36, 48, 60, 72, and 84 hours.

Cell fractionation and sodium borohydride treatment. Cell fractionation and borohydride treatment were done as previously reported (64) and are detailed in the Supplemental Methods.

Detection of serum mannose levels by GC-MS. A 100 μL serum sample was spiked with 1 μg $^{13}\text{C}_6$ -mannose as an internal standard, and 400 μL cold acetonitrile (-20°C) was added to precipitate the proteins. The sample was vortexed and kept on ice for 30 minutes to ensure

complete precipitation. The sample was then centrifuged at 14,000g at 4°C for 7 minutes, and the supernatant was removed and dried using a speed vac and lyophilizer. The dried samples were added to Tri-Sil Reagent (Thermo Fisher Scientific) and incubated at 80°C for 30 minutes. The reaction tube was then cooled to room temperature (RT), and excess reagent was removed by drying the samples. The Tri-Sil-derivatized sample was then dissolved in hexane and analyzed by GC-MS. Quantification was done by comparing the intensities of $^{12}\text{C}_6$ and $^{13}\text{C}_6$ mannose in the samples.

Detection of mannose in urine samples by GC-MS. Urine samples were centrifuged at 14,000g for 5 minutes at 10°C to remove any insoluble material. A 1 mL urine sample was spiked with 1 μg $^{13}\text{C}_6$ -mannose as an internal standard and passed over preconditioned LC-NH2 SPE tubes (Supelco), and the flow-through was collected. The LC-NH2 SPE tubes were further washed with 1 mL ultrapure water, and the dried flow-through products were then re-dissolved in 1 mL water, transferred into a screw-capped glass reaction tube, and dried in a lyophilizer. Tri-Sil HTP Reagent (250 μL , Thermo Fisher Scientific) was added on the dried sample, sonicated (10–15 s), and reacted at 80°C for 30 minutes. Tri-Sil-derivatized samples were then extracted by adding 1.5 mL hexane followed by sonication, vortexing, and filtration using a glass wool fiber-packed glass Pasteur pipette. The filtrate was dried and placed in the GC-MS autosampler vial. GC-MS analysis was performed in EI mode using the Restek-5MS glass capillary column (30 mL, 25 mm inner diameter [ID]). Identification of mannose was done from the retention time of mannose on the column and quantified by comparing the ratio of ion intensities (m/z of 204:206) of the corresponding fragment mass from $^{12}\text{C}_6$ and $^{13}\text{C}_6$ mannose.

Detection of serum/urine Sia levels by HPLC. Serum samples (250 μL) were spun at 20,000g for 10 minutes and filtered, and the DMB-derivatized samples were then analyzed by HPLC as previously described (97). Cold acetonitrile (400 μL , -20°C) was added to a 100 μL urine sample and kept on ice for 30 minutes. The sample was then centrifuged at 14,000g at 4°C for 7 minutes, and the supernatant was passed through a 3K spin filter. The flow-through was dried, DMB-tagged, and detected using HPLC-fluorescence detector (HPLC-FL) under isocratic solvent conditions (same as for Sia tagging).

Flow cytometry. Cells were blocked with 1% BSA in PBS and stained with mouse anti-Kdn antibodies on ice for 30 minutes. Following primary antibody staining, the cells were washed and stained with APC-conjugated anti-mouse antibody (Poly4053, BioLegend). Anti-Flag-F3165 (MilliporeSigma) was used to measure the population of *Cmas*-transfected HEK293A cells. Acquisition and analysis were performed using the BD FACSCalibur and FlowJo software, respectively.

Sequence alignments, tests for selection, and the phylogenetic tree. Coding nt and aa sequences of mannose metabolism genes and proteins were obtained from the NCBI's nt and protein databases. Nucleotide alignments and the percentage of identity were produced using the NCBI's Basic Local Alignment Search Tool (BLAST) (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). Protein sequences were aligned using the Clustal Omega program (UniProt.org). Nucleotide alignments were used to analyze the ratio of nonsynonymous (K_a) to synonymous (K_s) substitution, evaluated with an online calculation tool (<http://services.cbu.uib.no/tools/kaks>). The K_a/K_s ratio is representative of purifying selection on a coding sequence, as a K_a/K_s of less than 1 indicates that there is pressure to conserve the amino sequence relative to the background mutation rate. A maximum likelihood tree was constructed

using the general time reversible model (98). The tree with the highest log likelihood is shown. Initial trees for the heuristic search were obtained automatically by applying the maximum parsimony method. A discrete γ distribution was used to model evolutionary rate differences among sites (5 categories, +gamma distribution [+G], parameter = 0.6749). The rate variation model allowed for some sites to be evolutionarily invariable ([+invariance] (+I), 21.44% sites).

Detection of anti-Sia antibodies in pooled human intravenous Ig. Intravenous Ig (IVIg) was used for the detection of anti-Sia antibodies (Kdn, Neu5Gc, Neu5Ac). Briefly, a 96-well plate (CoStar) was coated overnight with 100 $\mu\text{g}/\text{mL}$ streptavidin at 4°C and washed with PBS containing 0.1% Tween 20 (PBST). The chemoenzymatically synthesized, defined glycans were used to determine the presence of antibodies against a specific glycan structure. The glycan epitopes used were Neu5Ac α 2-3Gal β 1-4GlcNAc-, Neu5Gc α 2-3Gal β 1-4GlcNAc-, Kdn α 2-3Gal β 1-4GlcNAc-, and unsialylated Gal β 1-4GlcNAc- (negative control). Each of these glycans was biotinylated. Following streptavidin coating, the wells were further blocked with PBST for 1.5 hours at RT and coated with 500 ng of the respective biotinylated glycan per well for 2 hours. Following incubation, the wells were washed with PBST and incubated with IVIg (1:250 dilution) for 1.5 hours at RT. Finally, the wells were washed and incubated with HRP-conjugated goat anti-human IgG antibodies (catalog 172-1050, Bio-Rad, 1:10,000 dilution) for 1 hour, and then washed and visualized at 490 nm following HRP substrate addition. For quantification of intensity resulting from the specific antibodies against Neu5Ac/Neu5Gc/Kdn epitopes and not from any nonspecific antibodies against the underlying glycan structure, the absorbance of the unsialylated conditions (wells coated with Gal β 1-4GlcNAc- glycans) was subtracted from the corresponding values of the sialoglycan-coated wells.

Statistics. Data were analyzed by *t* test or ANOVA, as indicated in the figure legends, and are presented as the mean (SD) using GraphPad Prism software, version 9 (GraphPad Software). *P* values of less than 0.05 were considered significant.

Study approval. Human sample collection was approved by the UCSD Human Research Protections Program (HRPP) (111279XX). Informed consent was obtained from all participants prior to sample collection. Ethics approval for the human mannose ingestion study was obtained from the UCSD HRPP (172014X).

Author contributions

AV conceptualized the study. K. Kawanishi, SS, SD, AS, BC, AMK, and AV designed the study methodology. SS and MV validated the findings. K. Kawanishi, SD, MV, AS, SSS, BC, and AMK conducted formal analysis. K. Kawanishi, SS, SD, MV, AS, SSS, BC, and AMK performed experiments. ICS, CS, K. Kitajima, KS, HHF, AMK, and AV provided resources. K. Kawanishi and AV wrote the original draft of the manuscript. K. Kawanishi, SS, SD, MV, XC, HHF, AMK, and AV reviewed and edited the manuscript. K. Kawanishi, SS, MV, SSS, AS, and AMK performed visualization of the data. HHF, AMK, and AV supervised the study. AV was responsible for project administration. HHF, and AV acquired funding for the study. K. Kawanishi designed study methodology, performed experiments, visualization of the data, and formal analysis, and wrote the original draft and reviewed and edited the manuscript. SS designed study methodology and performed data validation, experiments, visualization of the data, and reviewed and edited the manuscript.

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