The Biomarker GlycA Is Associated with Chronic Inflammation and Predicts Long-Term Risk of Severe Infection

Highlights

- Elevated GlycA was stable within individuals for up to a decade
- GlycA marked the levels of myriad inflammatory cytokines in circulation
- A gene network enriched for neutrophil functions was associated with GlycA
- GlycA strongly predicted future risk of hospitalization and death from infection

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In Brief
Ritchie et al. investigate the biology of GlycA, a known biomarker for short-term mortality. They reveal GlycA’s long-term behavior in apparently healthy patients: it is stable for >10 years and associated with chronic low-grade inflammation. Accordingly, GlycA predicts death from infection up to 14 years in the future.
The biomarker glycoprotein acetylation (GlycA) has been shown to predict risk of cardiovascular disease and all-cause mortality. Here, we characterize biological processes associated with GlycA by leveraging population-based omics data and health records from >10,000 individuals. Our analyses show that GlycA levels are chronic within individuals for up to a decade. In apparently healthy individuals, elevated GlycA corresponded to elevation of myriad inflammatory cytokines, as well as a gene coexpression network indicative of increased neutrophil activity, suggesting that individuals with high GlycA may be in a state of chronic inflammatory response. Accordingly, analysis of infection-related hospitalization and death records showed that increased GlycA increased long-term risk of severe non-localized and respiratory infections, particularly septicemia and pneumonia. In total, our work demonstrates that GlycA is a biomarker for chronic inflammation, neutrophil activity, and risk of future severe infection. It also illustrates the utility of leveraging multi-layered omics data and health records to elucidate the molecular and cellular processes associated with biomarkers.

INTRODUCTION

The integration of large-scale, systems-wide biomolecular information with health records to identify statistical associations represents a foundation of future studies in "precision medicine." Early systems-wide studies have elucidated the etiology of complex diseases in natural human populations (Chen et al., 2012, 2008; Emilsson et al., 2008; Montoya et al., 2014; Stanberry et al., 2013; Wang et al., 2011; Zhang et al., 2013). In doing so, they have identified biomarkers of potential clinical utility. Exemplar are recent studies that used nuclear magnetic resonance (NMR) spectroscopy biomarker profiling to identify serum glycoprotein acetylation (GlycA) levels as a biomarker for risk of 5-year all-cause mortality and incident cardiovascular disease (CVD) events (Akinkuolle et al., 2014; Fischer et al., 2014). Fischer et al. showed that elevation of GlycA was predictive of death from all causes within 5 years in >17,000 generally healthy individuals from two independent population-based cohorts (Fischer et al., 2014). Each SD increase of GlycA above the population mean conferred a 67% and 55% increase in mortality risk in the 5-year follow-up period for Estonian and Finnish cohorts, respectively (Fischer et al., 2014). Similarly, Akinkuolle
et al. investigated GlycA in a recent study of 27,500 initially healthy women. They found that elevated GlycA was predictive of 15-year risk of CVD incidence and mortality (Akinkuolie et al., 2014). In both studies, the predictive capacity of GlycA was independent of age, sex, modifiable lifestyle risk factors, medication, and disease prevalence. Other studies on GlycA in relation to common chronic diseases are rapidly emerging (Akinkuolie et al., 2015; Ala-Korpela, 2015); however, the biological foundation of these associations remains unexplored.

GlycA itself is a complex heterogeneous NMR signal that reflects the abundance of mobile N-acetyl sugar groups found on glycoproteins in circulating blood (Bell et al., 1987; Otvos et al., 2015). Multiple circulating glycoproteins have been found to contribute to GlycA, including alpha-1-acid glycoprotein, alpha-1 antitrypsin, haptoglobin, transferrin (Bell et al., 1987), and, more recently, alpha-1 antichymotrypsin; however, their relative contribution to the GlycA signal remains unclear (Otvos et al., 2015). The glycoproteins contributing to GlycA are involved in the so-called “acute-phase response” to exogenous insults, including infection and physical injury. This is a broad set of systemic biochemical and physiological changes, primarily driven by cytokine production of inflammatory cells, which occur immediately following the insult (Gabay and Kushner, 1999). With the exception of transferrin, the circulating concentrations of the proteins that constitute the GlycA signal are known to increase during the acute-phase response (Aronsen et al., 1972; Gitlin and Colten, 1987). Accordingly, the GlycA signal itself is elevated during the acute phase in patient samples (Bell et al., 1987).

Although GlycA is understood to mark the acute-phase response, it has also been shown to be elevated in patients with a diverse range of inflammation-linked chronic conditions, including rheumatoid arthritis, hypertension, obesity, and metabolic disorders (Bell et al., 1987; Lauridsen et al., 2010; Otvos et al., 2015; Würzt et al., 2012, 2014). Clinical assessment of chronic inflammation is routinely used in patient monitoring, disease diagnosis, and risk assessment. Currently, the state of the art for assessing chronic inflammation in the clinic typically includes quantification of circulating C-reactive protein (CRP), a single molecular species for which highly sensitive immunoassays are widely available (Pearson et al., 2003). However, the efficacy of this single marker may be limited by false-negatives. Chronic inflammation is likely as heterogeneous as the acute-phase response, where biochemical and physiological changes are not uniform across inflammatory conditions (Ebersole and Cappelli, 2000; Morley and Kushner, 1982). Evidence is now emerging to suggest that elevated CRP is not a universal feature of chronic inflammation. Importantly, GlycA, a composite marker of multiple molecular species, has been shown to predict risk of all-cause mortality and CVD, even when adjusting for CRP (Akinkuolie et al., 2014; Fischer et al., 2014).

This study elucidates the molecular and cellular processes underlying circulating GlycA levels by utilizing a range of omics data and electronic health records from three large-scale population-based cohorts (n = 11,825 total). Our unbiased integrative analysis reveals that GlycA levels are persistent over time and positively associated with both diverse cytokines in individuals without recent infection and the neutrophil antimicrobial response. This suggests that apparently healthy individuals may be in a state of chronic inflammatory response. Accordingly, we demonstrate that GlycA is predictive of long-term risk of severe infection using a large prospective cohort with nearly 14 years of follow-up health records. The overall study flow is illustrated in Figure 1.

**RESULTS**

**Study Data**

In this study, we analyzed data collected by three population-based studies (Supplemental Experimental Procedures). The Dietary, Lifestyle, and Genetic Determinants of Obesity and Metabolic syndrome (DILGOM) Study is a cross-sectional cohort of 579 individuals (300 female; 52%) 25–74 years old (Inouye et al., 2010a, 2010b). The 1997 collection of the FINRISK Study is a cross-sectional cohort of 7,599 individuals (3,822 female; 50%) 24–74 years old with disease events followed prospectively via nation-wide electronic health records (Borodulin et al., 2015; Würzt et al., 2015). The Cardiovascular Risk in Young Finns Study (YFS) is a longitudinal population-based study of 3,596 individuals (1,832 female; 51%) recruited during childhood in 1980 (Raitakari et al., 2008). In this study, data were available from the 2001, 2007, and 2011 follow-up collections (YFS2001, YFS2007, and YFS2011, respectively) (Würzt et al., 2014). By the 2011 follow-up, individuals were 34–49 years old. Clinical and demographic characteristics for each study are summarized in Table 1.

For all studies, NMR biomarker profiling data, including data for GlycA, were available (Soininen et al., 2009), as were CRP immunoassays. To fine-map the GlycA signal, alpha-1-acid glycoprotein, alpha-1 antitrypsin, haptoglobin, and transferrin...
were quantified by immunoturbidimetric methods in the DILGOM Study. Multiplexed cytokine panels were performed for YFS2007. Cytokine characteristics are reported in Table S1.

Fasting whole-blood transcriptomic data (Illumina HT-12 arrays) were available for the DILGOM Study and YFS2011. Electronic health records were collated for the FINRISK Study only. There was an insufficient number of endpoint events for GlycA survival analysis in the DILGOM Study due to small sample size and short follow-up time. Because of their relative youth and short follow-up time, there were also insufficient numbers of endpoint events in the YFS. Unless otherwise noted, all molecular measurements in all cohorts were performed on fasting serum samples. Protocols and processing details for each platform and study are described in the Supplemental Experimental Procedures.

### Specific Glycoprotein Levels Contributing to GlycA in a Population-Based Study

In contrast to previous studies that used protein standards to deconvolute the GlycA signal (Bell et al., 1987; Otvos et al., 2015), we utilized a population-based approach through the DILGOM Study to determine the relative contributions of serum alpha-1-acid glycoprotein, alpha-1 antitrypsin, haptoglobin, and transferrin levels to GlycA (Supplemental Experimental Procedures). As expected, we found that all four glycoproteins significantly contributed to GlycA signal intensity. Univariable linear regression indicated that each SD increase of alpha-1-acid glycoprotein, haptoglobin, alpha-1 antitrypsin, and transferrin corresponded to a 0.54 (p = 2 × 10^-3), 0.52 (p = 9 × 10^-4), 0.29 (p = 3 × 10^-2), and 0.20 (p = 1 × 10^-2) SD increase of GlycA.

### Table 1. Cohort Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>YFS 2001</th>
<th>YFS 2007</th>
<th>YFS 2011</th>
<th>DILGOM</th>
<th>FINRISK</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of individuals</td>
<td>2,232</td>
<td>2,159</td>
<td>2,041</td>
<td>579</td>
<td>7,599</td>
</tr>
<tr>
<td>Number (and %) of women</td>
<td>1,234 (55%)</td>
<td>1,185 (55%)</td>
<td>1,112 (54%)</td>
<td>300 (52%)</td>
<td>3,822 (50%)</td>
</tr>
<tr>
<td>Mean age in years (and range)</td>
<td>31.7 (24–39)</td>
<td>37.7 (30–45)</td>
<td>41.9 (34–49)</td>
<td>52.1 (25–74)</td>
<td>48.3 (24–74)</td>
</tr>
<tr>
<td>Body mass index (kg/m²)</td>
<td>25.1 ± 4.4</td>
<td>26.0 ± 4.8</td>
<td>26.5 ± 5.1</td>
<td>26.8 ± 4.7</td>
<td>26.7 ± 4.5</td>
</tr>
<tr>
<td>GlycA (mmol/l)</td>
<td>1.4 ± 0.2</td>
<td>1.6 ± 0.3</td>
<td>1.6 ± 0.2</td>
<td>1.6 ± 0.2</td>
<td>1.4 ± 0.2</td>
</tr>
<tr>
<td>Triglycerides (mmol/l)</td>
<td>1.3 ± 0.8</td>
<td>1.4 ± 0.9</td>
<td>1.3 ± 0.8</td>
<td>1.3 ± 0.6</td>
<td>1.5 ± 1.1</td>
</tr>
<tr>
<td>CRP (mg/l)</td>
<td>1.9 ± 3.8</td>
<td>1.9 ± 3.8</td>
<td>1.7 ± 3.2</td>
<td>2.7 ± 5.7</td>
<td>2.5 ± 5.3</td>
</tr>
<tr>
<td>Number of recent febrile infection</td>
<td>112 (5.0%)</td>
<td>95 (4.4%)</td>
<td>93 (4.6%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Use of antihypertensive therapy</td>
<td>7 (0.3%)</td>
<td>45 (2.1%)</td>
<td>74 (3.6%)</td>
<td>86 (15%)</td>
<td>269 (0.4%)</td>
</tr>
<tr>
<td>Prevalent diabetes</td>
<td>–</td>
<td>–</td>
<td>35 (1.3%)</td>
<td>61 (10%)</td>
<td>440 (5.8%)</td>
</tr>
<tr>
<td>Incident diabetes</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>8 (1.4%)</td>
<td>605 (8.0%)</td>
</tr>
<tr>
<td>Prevalent CVD</td>
<td>–</td>
<td>–</td>
<td>20 (0.7%)</td>
<td>17 (2.9%)</td>
<td>263 (3.5%)</td>
</tr>
<tr>
<td>Incident CVD</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>7 (1.2%)</td>
<td>802 (11%)</td>
</tr>
<tr>
<td>Prevalent cancer</td>
<td>–</td>
<td>–</td>
<td>40 (1.4%)</td>
<td>18 (3.1%)</td>
<td>180 (2.4%)</td>
</tr>
<tr>
<td>Incident cancer</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>11 (1.9%)</td>
<td>12 (0.2%)</td>
</tr>
<tr>
<td>Prevalent immunodeficiency</td>
<td>–</td>
<td>–</td>
<td>10 (0.4%)</td>
<td>0 (0%)</td>
<td>8 (0.1%)</td>
</tr>
<tr>
<td>Incident immunodeficiency</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0 (0%)</td>
<td>12 (0.2%)</td>
</tr>
<tr>
<td>Total number of deaths</td>
<td>–</td>
<td>–</td>
<td>15 (2.6%)</td>
<td>839 (11%)</td>
<td></td>
</tr>
<tr>
<td>Number of glycoprotein assays</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>579</td>
<td>–</td>
</tr>
<tr>
<td>Alpha-1-acid glycoprotein (g/l)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.8 ± 0.2</td>
<td>–</td>
</tr>
<tr>
<td>Alpha-1 antitrypsin (g/l)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1.2 ± 0.2</td>
<td>–</td>
</tr>
<tr>
<td>Haptoglobin (g/l)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1.1 ± 0.5</td>
<td>–</td>
</tr>
<tr>
<td>Transferrin (g/l)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>2.7 ± 0.4</td>
<td>–</td>
</tr>
<tr>
<td>Number of cytokine assays</td>
<td>–</td>
<td>2,159</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<tr>
<td>Cell count data</td>
<td>–</td>
<td>–</td>
<td>2,021</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Gene expression profiling number</td>
<td>–</td>
<td>–</td>
<td>1,649</td>
<td>518</td>
<td>–</td>
</tr>
</tbody>
</table>

Data are reported as the mean ± SD unless otherwise indicated. Prevalent disease indicates events occurring prior to sample collection, while incident disease indicates events occurring after sample collection. For the YFS, a disease was considered prevalent if it occurred any time prior to the 2011 sample collection for any of the 2,766 individuals participating at any one time point. Cytokine characteristics are reported in Table S1. Dashes indicate that data are not applicable.
respectively. Of the individuals in the top 10% for the GlycA levels, 40% were also in the top decile for alpha-1-acid glycoprotein levels, as were 43% for haptoglobin, 29% for alpha-1 antitrypsin, and 15% for transferrin. Pairwise correlations among the glycoproteins were either positive or, in the case of alpha-1-acid glycoprotein and transferrin, not significant. These analyses indicate that the GlycA signal is not dominated by any single glycoprotein; thus, GlycA is treated as a composite signal in all subsequent analyses.

GlycA Levels in Acute Phase, Chronicity over Time, and Relationship with Myriad Cytokines

All YFS collections contained samples from individuals reporting febrile infection within 2 weeks prior to sample collection. In these individuals, we observed modest elevation of GlycA in individuals from all YFS collections (mean increase of 0.41 SD; Figure S1), consistent with GlycA’s role as a positive acute-phase reactant. Among YFS individuals not reporting a recent febrile infection, there was a strong autocorrelation of GlycA levels within individuals across all three time points (mean Pearson r = 0.43). Of the top 10% of individuals by GlycA level in 2001, 20% were also in the top decile in 2007, and 19% were also in the top decile in 2011. Individuals reporting a recent febrile infection were no more likely than others to have elevated GlycA levels at any subsequent time point.

Given that elevated GlycA levels correlated with both the acute-phase response and were stable over time, we asked whether GlycA was associated with inflammatory signatures in apparently healthy individuals. We utilized multiplex cytokine panels assayed in YFS2007 serum samples (Supplemental Experimental Procedures) from individuals reporting no recent febrile infections and observed a large number of robust associations (Bonferroni-adjusted p < 0.001): 30 of 36 tested cytokines were significantly associated with GlycA (Figure 2). The concentrations of 29 cytokines, with both anti- and pro-inflammatory roles, increased along with the GlycA signal (Figure 2). Only cutaneous T-cell-attracting chemokine (CTACK) showed a significant negative association with GlycA (Figure 2). Cytokine concentration changes associated with GlycA were relatively modest, with even the top decile of GlycA levels corresponding to cytokine elevations of 0.11–0.48 SD on average (Figure 2). Together, these results suggest an association between elevated GlycA and systemic low-grade inflammation in an apparently healthy population. It was not possible to test whether cytokine concentrations were stable over time, like those of GlycA, because cytokine panels were not available for YFS2001 or YFS2011.

A Transcriptional Subnetwork Associated with GlycA

When put in the context of previous studies, our results suggest that, in addition to marking the acute-phase response (Bell et al., 1987), high GlycA levels may also mark chronic inflammation in apparently healthy individuals. This suggests an intimate connection between the circulating, systemic concentration of GlycA and cellular-level changes in physiology. To gain insight into cellular processes associated with elevated GlycA, we utilized whole-blood transcriptome profiling in the DILGOM and YFS2011 cohorts. We inferred weighted gene coexpression network modules (Zhang and Horvath, 2005) in the DILGOM cohort and then tested each module for an association with the GlycA signal (Supplemental Experimental Procedures). Briefly, the full gene coexpression network was inferred as the pairwise Spearman correlation between all microarray probes. This was converted to a measure of network adjacency through a soft-threshold power transform. Modules were then detected through a dynamic tree cut algorithm on the topological overlap similarity of probe adjacencies. A summary expression vector (first principal component) was calculated for each module and used to test for the association with GlycA. Models were adjusted for age, sex, and, additionally, serum triglyceride levels, which have been shown to modify the multivariable effect of lipoprotein concentrations on the GlycA signal and associated mortality risk (Fischer et al., 2014). Significantly associated modules were subsequently tested for replication in the YFS cohort using NetRep (v0.10.4), a rapid permutation-based approach of
We identified 40 coexpression modules within the DILGOM whole-blood expression data, of which 3 were significantly associated with GlycA after adjusting for multiple testing ($p < 0.001$). One module’s network topology and GlycA association was replicated in the YFS2011 whole-blood expression data (permutation test $p$ value $< 0.01$ for all network-based statistics and GlycA association $p$ value $< 0.001$; Table S2). We subsequently performed in-depth analysis of the module (Figure 3) and its 27 genes (Table S3). For this module, single-gene associations with GlycA were consistent with the whole-module association with GlycA.

The enriched GO processes and KEGG pathways of the module largely implicated innate immune response to microbes (Table S4). Consistent with this, module expression was elevated by 0.61 SDs ($p = 3 \times 10^{-4}$) in individuals reporting recent febrile infection (Figure 3). Most genes in the preserved module were primarily reported in the literature as coding for antimicrobial peptides synthesized in and secreted from neutrophil granules (14 genes, 52%; Figure 3; Table S5). Genes with antimicrobial function constituted 11 of the top 14 genes when ranked by mean intramodular connectivity across datasets (Table S3), a topological measure known to correspond to relative biological importance (Langfelder et al., 2013). A further three genes coded for proteins serving a non-antimicrobial role in neutrophil-granule-associated processes (Table S5), and two genes coded for proteins expressed on the surface of neutrophils (Table S5). Since a majority of these genes have well-characterized antimicrobial roles in neutrophil granules (Faurusch and Borregaard, 2003), we refer hereinafter to this module as the neutrophil module.

Neutrophils are the dominant white blood cell type in adult humans, but their abundance varies in populations and within individuals over time (Summers et al., 2010). While neutrophil cell counts were not available, we observed a strong positive association between overall leukocyte abundance and elevated neutrophil module summary expression ($b = 0.24$ SD leukocyte abundance increase per SD increase of neutrophil module expression, $p = 2 \times 10^{-6}$), suggesting that elevated neutrophil module expression may be associated, to some extent, with increased circulating neutrophil abundance. Adjustment of the GlycA model for leukocyte abundance showed that neutrophil module expression retained a strong association with GlycA; however, both the module and leukocyte abundance were highly significant ($b = 0.09$, $p = 2 \times 10^{-6}$, and $b = 0.15$, $p = 2 \times 10^{-15}$, respectively), indicating that increased GlycA corresponds to increased production of both antimicrobial peptides and circulating leukocytes.

Figure 3. Neutrophil Gene Coexpression Module Associated with GlycA

(A and B) Probe coexpression (Spearman correlation) in the DILGOM cohort (A) and replication in the YFS2011 cohort (B). See Table S3 for module gene details. Symbols above gene symbol indicate expression in neutrophils (#), whether its product is located in neutrophil granules (%), and whether its product has antimicrobial functionality (*). See Table S5 for a summary of the literature for each gene.

(C) Replication of GlycA association in both cohorts. Associations were assessed using a linear regression of GlycA on the module summary expression adjusting for age, sex, and triglycerides. Magnitude corresponds to change in SD units of GlycA per SD increase of neutrophil module expression. CI, confidence interval.

(D) The top enriched GO biological process terms. A full list of significant GO terms and KEGG pathway enrichments can be found in Table S4.

(E) Association of neutrophil module expression with alpha-1-acid glycoprotein, haptoglobin, alpha-1 antitrypsin, and transferrin in the DILGOM cohort in a multivariable linear regression. Magnitude corresponds to change in SD units of GlycA per SD increase of each respective acute-phase glycoprotein.

(F) Boxplots showing association with recent febrile infection (self-reported) ($n = 72$ with febrile infection, and $n = 1,550$ without). T-test difference, 95% confidence interval, and $p$ value are reported. GlycA and triglyceride levels were log-transformed. All continuous measurements were standardized.

See also Tables S3, S4, and S5.

suggesting an overabundance of neutrophils and that their granule content may directly contribute to elevated GlycA.

**GlycA Is Associated with Risk of Hospitalization and Death from Microbial Infection**

Because GlycA and the microbial-associated inflammatory response were statistically associated, and it is known that the inflammatory response correlates with adverse outcomes in the clinic, we hypothesized that GlycA level may also be reflective of future infection severity risk. We asked whether there was a relationship between risk of future hospitalization or death and elevated GlycA levels. To do this, we assessed infection-related causes of hospitalization or death in a population-based collection of 7,599 individuals from FINRISK 1997 who had baseline GlycA profiles and electronic health records with 13.8 years of follow-up (Supplemental Experimental Procedures). Cox models were fitted to each infectious disease event and all-cause mortality (Akinkuolie et al., 2014; Fischer et al., 2015), which are, themselves, responsive to myriad inflammatory stimuli (Gabay and Kushner, 1999). Previous studies have shown that GlycA is predictive of hospitalization from infection, including fungal infections and localized infections of the skin, bones, joints, intestinal, and urinary system (Figure 4B). Adjustment of GlycA models for prevalent chronic diseases, compared to CRP, made little difference to GlycA HRs for future risk of hospitalization and death from infection (Figure S3A). GlycA displayed stronger prediction on average than CRP across all infection-related events (Figure S3B), and adjustment of the models for CRP only slightly attenuated the GlycA HRs (Figure S3C). Taken together, these results suggest that apparently healthy individuals with elevated baseline levels of GlycA are at increased risk for severe infection events for up to 14 years in the future.

**DISCUSSION**

GlycA is a composite NMR-based signal related to changes in multiple circulating glycoproteins (Bell et al., 1987; Otvos et al., 2015), which are, themselves, responsive to myriad inflammatory stimuli (Gabay and Kushner, 1999). Previous studies have shown that GlycA is predictive of long-term risk for CVD events and all-cause mortality (Akinkuolie et al., 2014; Fischer et al., 2014). In this study, we characterized the molecular and cellular processes underlying GlycA variation in circulation. In apparently healthy individuals, we have shown that GlycA can be chronically elevated for periods of up to a decade; in individuals with high GlycA, widespread, modest elevation of numerous cytokines are suggestive of a prolonged low-grade inflammatory state. Accordingly, we identified a robust transcriptional module in whole blood that was strongly positively correlated with baseline GlycA levels, suggesting an overabundance of neutrophils and that their granule content may directly contribute to elevated GlycA.
associated with GlycA levels and enriched for innate immune response genes—specifically, those with neutrophil antimicrobial function. Elevated GlycA levels had long-term ramifications for risk of infection: high GlycA levels correlated with an increased risk of hospitalization and death from diverse infections—particularly septicemia and pneumonia—that persists for over a decade.

The question remains as to the cause of GlycA-associated chronic inflammation. Taken together, our results suggest multiple potential scenarios where apparently healthy individuals with high GlycA levels have (1) persistent but clinically silent, low-grade microbial infection(s), (2) a past severe infection event that results in modest chronic inflammation, (3) low-grade, chronic inflammation that mimics an anti-microbial response, or (4) a combination of these. This GlycA-related immune response was characterized by a gene coexpression network that harbored a clear neutrophil transcriptional signature indicative of antimicrobial function (Faurschou and Borregaard, 2003). Once recruited to the site of infection, neutrophils secrete antimicrobial peptides from their granules into the extracellular matrix and into phagosomes that surround microbes. They also form neutrophil extracellular traps (NETs): extracellular fibers composed of neutrophil DNA and granule antimicrobial peptides (Brinkmann et al., 2004; Urban et al., 2009). Neutrophil antimicrobial peptides are generally cytotoxic, damaging and degrading proximal tissue. aberrantly increased neutrophil activity has been observed in a range of chronic inflammatory conditions that are generally thought to be independent of microbial infection, including rheumatoid arthritis (Branz and Papayannopoulos, 2013; Wright et al., 2010), and increased neutrophil abundance itself has been shown to have predictive capacity for long-term mortality and myocardial infarction event risk (Horne et al., 2005).

Consistent with the diverse ramifications of neutrophil-driven inflammation, Warnatsch et al. recently reported a causal link between cholesterol, NET formation, and pro-inflammatory response driving atherosclerotic plaque growth (Warnatsch et al., 2015). Notably, two of the genes we identify as part of the neutrophil module, ELANE and MPO, have key roles in NET formation (Papayannopoulos et al., 2010). Furthermore, the long-term risk of severe infection events in apparently healthy individuals with elevated GlycA levels was consistent with overactivity of immune response, though GlycA may not necessarily play a causal role. Overactivity of immune response from severe infection events, such as sepsis and influenza, is known to cause tissue damage and organ dysfunction through hypercytokinemia (Tisoncik et al., 2012), and baseline elevation of circulating cytokines correlated with GlycA may contribute to its association with hospitalization and mortality risk. While our findings are largely observational, the prospect of systematic integration of prior medical interventions may yield causal relationships. In addition, an immediately fruitful avenue of research may be in vivo assays of neutrophil function, using neutrophils derived from individuals with variable GlycA levels.

This study demonstrates the utility of an approach that integrates diverse omics data and long-term follow-up of health records to uncover putative cellular-level processes associated with a recently uncovered biomarker. As multi-omics studies become increasingly common, biomarker associations with medical histories and health outcomes are likely to improve both patient care and our knowledge of the basic biological mechanisms underlying human disease. This study represents a potentially useful strategy for leveraging omics information for future studies in precision medicine.

**EXPERIMENTAL PROCEDURES**

Full experimental procedures can be found in the Supplemental Experimental Procedures. For YFS, ethics were approved by the Joint Commission on Ethics of the Turku University and the Turku University Central Hospital; for DILGOM, ethics were approved by the Coordinating Ethical Committee of the Helsinki and Uusimaa Hospital District; and for FINRISK, ethics were approved by the ethical committee of the National Public Health Institute, Finland (Supplemental Experimental Procedures).

**Data Collection and Quantification**

Concentrations of GlycA and triglycerides were quantified from serum samples using a proton NMR platform (Soininen et al., 2009). Concentrations of high-sensitivity CRP and the four glycoproteins were quantified from serum using immunonoturbidimetric assays, and the 36 cytokines were quantified using multiplex cytokine panels. Genome-wide gene expression profiling was carried out on whole blood using the Illumina HT12 platform, and blood cell count analysis was carried out using flow cytometry. Incident event data were obtained from the Finnish National Hospital Discharge Register and the National Causes of Death Register (Lim et al., 2014). Clinical characteristics were also obtained from survey questions. Data type availability, sample sizes, and cohort characteristics are provided in Table 1.

**Network Analysis**

Network inference and module detection were performed in the DILGOM cohort using weighted gene coexpression network analysis (WGCNA) (Zhang and Horvath, 2005). Network inference was adjusted for age and sex. The soft-threshold power used for network construction was 4 (Figure S4A). Repli- cation of the three GlycA-associated modules was assessed in the YFS2011 cohort through permutation testing of seven network-based statistics (Langfelder et al., 2011) (Table S6), using a pre-release of the NetRep R package (version 0.10.4, available at https://github.com/InouyeLab/NetRep/releases/tag/v0.10.4). For this purpose, the network inference component of WGCNA was performed on the YFS2011 gene expression. The soft threshold power was 4 (Figure S4b). Summary expression profiles were calculated as the eigenvector of the first principal component of each module’s gene expression matrix. We use the shorthand “neutrophil module expression” to refer to the neutrophil module’s summary expression profile throughout the text. Over-representation analysis of Gene Ontology (GO) terms and KEGG pathways was performed using the web tool, GeneCoDis (Garmona-Saez et al., 2007).

**Statistical Analyses**

A natural log-transform was applied to GlycA, each glycoprotein, triglycerides, and leukocyte abundance. A rank-based inverse normal transform was applied to each cytokine. Each measurement, including each module’s summary expression, was standardized to SD units. All regression models were adjusted for age and sex, and models testing association between GlycA and module expression were additionally adjusted for triglycerides. HRs of GlycA for hospitalization or death were assessed by Cox proportional-hazard models using age as the timescale and adjusted for sex, triglycerides, and prior occurrence of the diagnosis category within ten years prior to sample collection. Sensitivity analyses were performed by (1) additionally adjusting for prevalent chronic diseases or inflammatory disorders, (2) replacing GlycA for CRP and, (3) adjusting GlycA for CRP.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, and six tables and can be found with this article online at http://dx.doi.org/10.1016/j.cels.2015.09.007.
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