Co-occurrence of Type 1 Diabetes and Celiac Disease Autoimmunity

William Hagopian, MD, PhD; Hye-Seung Lee, PhD; Edwin Liu, MD; Marian Rewers, MD, MPH; Jin-Xiong She, PhD; Anette-G. Ziegler, PhD; Åke Lernmark, MD; Jorma Toppari, MD, PhD; Stephen S. Rich, PhD; Jeffrey P. Krischer, PhD; Henry Erlich, PhD; Beena Akolkar, PhD; Daniel Agardh, MD, PhD; the TEDDY Study Group

BACKGROUND AND OBJECTIVES: Few birth cohorts have prospectively followed development of type 1 diabetes (T1D) and celiac disease (CD) autoimmunities to determine timing, extent of co-occurrence, and associated genetic and demographic factors.

METHODS: In this prospective birth cohort study, 8676 children at high genetic risk of both diseases were enrolled and 5891 analyzed in median follow-up of 66 months. Along with demographic factors and HLA-DR-DQ, genotypes for HLA-DPB1 and 5 non-HLA loci conferring risk of both T1D and CD were analyzed.

RESULTS: Development of persistent islet autoantibodies (IAs) and tissue transglutaminase autoantibodies (tTGAs), as well as each clinical disease, was evaluated quarterly from 3 to 48 months of age and semiannually thereafter. IAs alone appeared in 367, tTGAs alone in 808, and both in 90 children. Co-occurrence significantly exceeded the expected rate. IAs usually, but not always, appeared earlier than tTGAs. IAs preceding tTGAs was associated with increasing risk of tTGAs (hazard ratio [HR]: 1.48; 95% confidence interval [CI]: 1.15–1.91). After adjusting for country, sex, family history, and all other genetic loci, significantly greater co-occurrence was observed in children with a T1D family history (HR: 2.80), HLA-DR3/4 (HR: 1.94) and single-nucleotide polymorphism rs3184504 at SH2B3 (HR: 1.53). However, observed co-occurrence was not fully accounted for by all analyzed factors.

CONCLUSIONS: In early childhood, T1D autoimmunity usually precedes CD autoimmunity. Preceding IAs significantly increases the risk of subsequent tTGAs. Co-occurrence is greater than explained by demographic factors and extensive genetic risk loci, indicating that shared environmental or pathophysiological mechanisms may contribute to the increased risk.

WHAT’S KNOWN ON THIS SUBJECT: Children with type 1 diabetes (T1D) are at greater risk of celiac disease (CD), and screening them for transglutaminase autoantibodies (tTGAs) is useful. T1D and CD share high-risk HLA-DR-DQ genotypes DR3 and DR4, which partly explains disease overlap.

WHAT THIS STUDY ADDS: Early markers of T1D and CD autoimmunity coincide more than expected. Overlap associates with familial T1D, HLA-DR3/4, and SH2B3, but is not fully explained by known risk factors. T1D autoantibodies usually precede tTGAs and confer significant risk of subsequent tTGAs.

Celiac disease (CD) and Type 1 diabetes (T1D) involve autoimmune attacks on small bowel mucosa and on pancreatic islet β cells, respectively. Both diseases have a strong genetic predisposition and most often begin in early childhood. A subclinical or preclinical phase is characterized by autoantibody positivity with minimal symptoms. Later, more severe target organ dysfunction appears, prompting clinical diagnosis and treatment.

The frequent coexistence of T1D and CD is widely described, but most researchers measure future risk of CD in children with previous T1D. This ranged from 2.5% to 16.4% (5.7% overall) in 23 studies worldwide, much greater than background CD rates. Conversely, studies of subsequent T1D in children with previous CD also revealed significantly more T1D than in non-CD controls. Presence of either disease therefore appears to increase the risk of developing the other.

Commonly postulated reasons for concurrent CD and T1D include shared HLA genetic risk and shared environmental exposures. Genetics are indeed important in CD, with ~75% monozygotic twin concordance and 10% of first-degree relatives affected. Genetics are similarly important in T1D, with ~42% monozygotic twin concordance and 6% to 8% FDRs affected. These genetic effects are greater than for most autoimmune diseases, perhaps because of a remarkably large effect of HLA Class II on both diseases. HLA-DR-DQ accounts for ~40% to 50% of the overall T1D genetic risk, and 53% of that for CD, using modern prevalence estimates. The 2 common haplotypes conferring increased risk in both diseases are DRB1*04-DQA1*03:01-DQB1*03:02 (commonly referred to as DR3-DQ2 haplotype, abbreviated DR3). In T1D, DR4 confers greater risk than DR3, but DR3/DR4 heterozygotes are at the greatest risk. For CD, DR3 confers more risk than DR4, and DR3/DR3 confers the greatest risk. Studies of the absolute CD risk conferred by HLA-DR-DQ in unbiased populations (without symptoms, other autoimmunity, or family history [FH]) are sparse, but the Environmental Determinants of Diabetes in the Young (TEDDY) study recently published multicenter risk estimates for several high-risk HLA antigen genotypes that were consistent with most previous literature. HLA-DR-DQ strongly influences both diseases and the risk-conferring haplotypes are shared, so substantial disease overlap is expected solely on that basis. CD and T1D are also concurrently influenced by >10 other genetic loci, conferring smaller hazard ratios (HRs) in the 1.14 to 1.35 range but with independent effects that may be cumulative. However, for only 6 loci (HLA-DPB1, RGS1, SH2B3, CTLA4, CCR3/CCR5, and PTPN2) does the relevant polymorphism affect both diseases in the same direction, therefore potentially increasing disease co-occurrence.

The TEDDY study is an international study of 8676 infants with many years of frequent prospective observation. The major inclusion criterion of elevated HLA antigen risk for T1D also confers elevated CD risk. Frequent blood sampling enabled accurate timing of seroconversion for both islet autoantibodies (IAs) and tissue transglutaminase autoantibodies (tTGAs). The cohort underwent extensive additional genotyping. TEDDY thus represents an unprecedented opportunity to describe the extent and timing of co-occurrence of the 2 diseases. The aim of this study was to investigate if 1 disease might trigger the other, and to test if their co-occurrence can be explained by common inherited risk genes.

**METHODS**

**The TEDDY Cohort and HLA Genotyping**

TEDDY is a prospective cohort study designed to identify environmental causes of T1D. From 2004 to 2010, 424 788 newborns were screened at 6 US and European centers. TEDDY enrolled 8676 infants with high-T1D–risk HLA antigen genotypes by 4.5 months of age, with intent to manage until age 15 years. Eligible HLA-DR-DQ genotypes are abbreviated DR3/4, DR4/4, DR4/8, and DR3/3 (Supplemental Table 2). Of 8676 TEDDY enrollees, 5891 were analyzed herein on the basis of full autoantibody characterization, genotyping on the ImmunoChip, and carrying 1 of the 4 major TEDDY-eligible HLA genotypes. Frequencies of all eligible HLA genotypes for each center are given in Supplemental Table 3. At the time of analysis, median follow-up was 66 months (range 10–111 months, interquartile range [IQR] 50–81 months), covering 32 454 person-years of observation. Local institutional review board approval and parental informed consent was obtained for all children. The study is monitored by an External Evaluation Committee of the US National Institutes of Health.

**Autoantibody Measurements**

Glutamate decarboxylase autoantibodies, insulinoma antigen-2 autoantibodies, and insulin autoantibodies (IAs) were measured in 2 harmonized core laboratories by using radiobinding assays incorporating extensive quality control. Persistent islet antibodies (IAs) were defined as positive antibodies to the same antigen confirmed by both core laboratories in 2 consecutive samples.
in 2 laboratories with final status determined by a central reference laboratory as previously described. Persistent tTGAs were defined as positive tTGAs test results in 2 consecutive samples. Only persistent IAs and tTGAs are considered in the current analyses.

**Prospective Follow-Up**

The TEDDY protocol specifies serum collection every 3 months from ages 3 to 48 months and at least every 6 months thereafter. IAs were measured in every available sample. Concurrent diseases including T1D and CD in the TEDDY child, and immediate family were noted at each visit. T1D was defined by using American Diabetes Association criteria. tTGA testing began at age 24 months and continued annually. If the results of any sample were positive, all previous samples from that child were analyzed to determine when tTGAs first appeared. Celiac symptoms were uncommon at the time of tTGA seroconversion. Parents of children with persistent tTGAs were notified and encouraged to pursue further evaluation under guidance of their physician. This likely led to more rapid ascertainment of CD in study than in general medical practice. In similar fashion, IA surveillance and early T1D ascertainment via glucose tolerance testing was also a feature of TEDDY. The decision on duodenal biopsy was outside of the study protocol, but if a biopsy were performed, a Marsh score of ≥2 defined CD. For individuals who did not receive a biopsy, a mean persistent tTGAs ≥100 U based on 2 consecutive samples was >95% specific for CD and therefore considered diagnostic herein. Children developing T1D were no longer on study protocol, but phone interviews and medical records reviews enabled determination of subsequent tTGA status, CD status, biopsy results if any, and use of a gluten-free diet in 150 out of 167 (90%) of these individuals. At the time of analysis, clinical T1D occurred in 138 (2.3%), CD in 323 (5.5%), and both diseases developed in 18 (0.3%) of the 5891 screened participants, (Fig 1). The low number of subjects concurrently diagnosed with T1D and CD in early childhood limited detailed subanalysis of factors influencing T1D and CD co-occurrence. We therefore focused on IAs and tTGAs as earlier and more prevalent disease markers.

**Single-Nucleotide Polymorphism Genotyping**

In addition to HLA-DR-DQ alleles, the cohort was genotyped for single-nucleotide polymorphisms (SNPs) by the Illumina ImmunoChip covering 186 genomic regions associated with 12 autoimmune diseases. SNP markers with call rates <90% or allele distributions strongly deviating from Hardy-Weinberg equilibrium in controls (P < 10^-6) were discarded. This resulted in data on ~176 586 SNPs in each of 7023 subjects. SNP genotypes were used to determine HLA-DPB1, and 5 other loci reported to affect both T1D and CD in the same direction: RGS1, SH2B3, CTLA4, CCR3/CCR5, and PTPN2 (Supplemental Table 4).

**Statistical Methods**

Excess co-occurrence was calculated by comparison with the expected along with the Wilson 95% confidence interval (CI). Excess proportion was compared by using Fisher’s exact test. To explore the pattern of appearance of each autoantibody (IAs and tTGAs) or disease (T1D and CD), the proportion of event-free survival in which Kaplan-Meier estimates were used was plotted against age of seroconversion or diagnosis. Age was compared between groups by using the Wilcoxon rank sum test.

A Cox proportional hazards model was used to investigate factors associated with IA and tTGA co-occurrence. Time to co-occurrence was defined as age of seroconversion.
The proportion of children who tested positive for IAs that developed T1D was 130 out of 457 (28.5%) compared with 8 out of 5434 (0.15%) of children who tested negative for IAs. The proportion of children who tested positive for tTGAs that developed CD was 321 out of 898 (35.8%) compared with 2 out of 4993 (0.04%) of children who tested negative for tTGAs. A total of 367 (6.2%) participants had IAs but not tTGAs, 808 (13.7%) had tTGAs but not IAs, and 90 (1.5%) had both (Fig 1). Assuming independence and assuming the observed proportion of tTGAs and IAs in TEDDY are accurate estimates of corresponding population proportions, the product of the observed IA and tTGA proportions represents the expected proportion of having both antibodies. The observed IA prevalence was 457 out of 5891 (7.8%), whereas that for tTGAs was 898 out of 5891 (15.2%). We therefore expected IAs and tTGAs to occur together in 1.19% (7.8%*15.2%) or 70 children. The observed prevalence of 90 out of 5891 (1.53%) revealed a significant excess of 20 children or 0.34% (95% CI: 0.21%–0.52%). Subgroups by country, sex, FH of T1D and of CD, HLA-DR-DQ, and other genetic loci each retained significant excess co-occurrence of IAs and tTGAs (Supplemental Table 5). In examining the proportion of children with each autoantibody, it can be seen that having IAs increased the proportion with tTGAs from 14.9% to 19.7% (Fig 2A). Conversely, having tTGAs increased the proportion with IAs from 7.4% to 10.0% (Fig 2B). This represents a 32% greater prevalence of tTGAs in those with IAs, and a 36% greater prevalence of IAs in those with tTGAs. Increases were observed across all individual HLA genotypes but were only significant for DR4/4 (P = .017) and for the combined data set of all HLA genotypes (P = .012).

Given the prospective nature of the TEDDY study, it is possible to extend the cross-sectional data of Fig 1 to examine the development of autoimmunity over time. Seroconversion for each of IAs and tTGAs was greatest in the first few years of life (Fig 3). When considered separately, the median age at seroconversion for IAs was 24 (IQR: 14–38.5) months and for tTGAs was 33 (IQR: 24–45) months (P < .0001). There was no significant difference in age at seroconversion when comparing those with 1 type of autoimmunity to those with both. Within the current follow-up, T1D was on average diagnosed earlier than CD (T1D at 36 [23–56] months; CD at 42 [34–57] months; P = .0007). Of the 90 individuals with both IAs and tTGAs, 5 (6%) developed the autoantibodies simultaneously, 61 (68%) developed IAs first at 24

For the following autoimmunity. The right-censored time was age at last sample collection. The model considered country, sex, and FH of T1D or CD (defined as an affected FDR), in addition to HLA-DR-DQ, HLA-DPB1, and the 5 other relevant genetic loci. Additionally, we examined the effect of each autoantibody on appearance of the other. For example, the effect of IAs preceding tTGAs on the risk of tTGAs was assessed (by using age of tTGA appearance as time to event) as compared with those without IAs at the time of tTGA appearance. Two sided P values < .05 were considered significant. SAS 9.4 Software (SAS Institute, Cary, NC) was used for all analyses.

**RESULTS**

The proportion of children who tested positive for IAs that developed T1D was 130 out of 457 (28.5%) compared with 8 out of 5434 (0.15%) of children who tested negative for IAs. The proportion of children who tested positive for tTGAs that developed CD was 321 out of 898 (35.8%) compared with 2 out of 4993 (0.04%) of children who tested negative for tTGAs. A total of 367 (6.2%) participants had IAs but not tTGAs, 808 (13.7%) had tTGAs but not IAs, and 90 (1.5%) had both (Fig 1). Assuming independence and assuming the observed proportion of tTGAs and IAs in TEDDY are accurate estimates of corresponding population proportions, the product of the observed IA and tTGA proportions represents the expected proportion of having both antibodies. The observed IA prevalence was 457 out of 5891 (7.8%), whereas that for tTGAs was 898 out of 5891 (15.2%). We therefore expected IAs and tTGAs to occur together in 1.19% (7.8%*15.2%) or 70 children. The observed prevalence of 90 out of 5891 (1.53%) revealed a significant excess of 20 children or 0.34% (95% CI: 0.21%–0.52%). Subgroups by country, sex, FH of T1D and of CD, HLA-DR-DQ, and other genetic loci each retained significant excess co-occurrence of IAs and tTGAs (Supplemental Table 5). In examining the proportion of children with each autoantibody, it can be seen that having IAs increased the proportion with tTGAs from 14.9% to 19.7% (Fig 2A). Conversely, having tTGAs increased the proportion with IAs from 7.4% to 10.0% (Fig 2B). This represents a 32% greater prevalence of tTGAs in those with IAs, and a 36% greater prevalence of IAs in those with tTGAs. Increases were observed across all individual HLA genotypes but were only significant for DR4/4 (P = .017) and for the combined data set of all HLA genotypes (P = .012).

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<table>
<thead>
<tr>
<th>HLA-DR-DQ Genotype</th>
<th>% of Cohort With IA</th>
<th>P = .017</th>
<th>% of Cohort With tTGA</th>
<th>P = .012</th>
</tr>
</thead>
<tbody>
<tr>
<td>DR3/3</td>
<td>35%</td>
<td></td>
<td>30%</td>
<td></td>
</tr>
<tr>
<td>DR3/4</td>
<td>30%</td>
<td></td>
<td>25%</td>
<td></td>
</tr>
<tr>
<td>DR4/4</td>
<td>25%</td>
<td></td>
<td>20%</td>
<td></td>
</tr>
<tr>
<td>DR4/8</td>
<td>20%</td>
<td></td>
<td>15%</td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>15%</td>
<td></td>
<td>10%</td>
<td></td>
</tr>
</tbody>
</table>

**FIGURE 2**

Cross-sectional analysis of autoantibody prevalence. A, Comparison of transglutaminase autoantibodies (CDA) prevalence in those with IAs versus those without IAs on fixed individual HLA-DR-DQ backgrounds as shown. B, Similar comparison of IA prevalence in those with or without tTGAs. The rightmost bars in each panel show the comparison when all 4 HLA antigen backgrounds are combined.
developed tTGAs first at 33 (24, 45) months (P < .0001). Comparative timing of IAs versus tTGAs was not different when comparing glutamate decarboxylase autoantibodies or IAAs as the specific IAs (Supplemental Fig 4). Greater HLA-DR3 gene dosage conferred shorter median time interval from development of IAs to tTGAs (median 2 months in DR3/3, 14 months in DR3/4, and 18 months in DR4/4 or DR4/8) (Supplemental Fig 5). The median time from tTGAs to CD was 11.6 (IQR: 7.9 – 16.1) months, significantly shorter than that from IAs to T1D at 17.2 (7.7 – 35.7) months (P < .0001).

A Cox Proportional Hazards model was used to examine risk factors for each individual type of autoimmunity, including whether each autoantibody influenced the risk for developing the other (Table 1). Known risk factors for each type of autoimmunity showed expected results in the adjusted model. Factors associated separately with increased IAs and with increased tTGAs included the countries of Finland or Sweden and HLA-DR3/4. HLA-DR3/3 increased tTGA risk but decreased IA risk versus the reference genotype. As expected, having an FDR with CD increased tTGA risk but not IA risk, and having an FDR with T1D increased IA risk but not tTGA risk. RG51 (rs2816316) and CTLA4 (rs3087243) were associated with increased TGA risk, whereas SH2B3 (rs3184504) was associated with increased IA risk. Importantly, in unadjusted analysis, IAs preceding tTGAs was associated with greater risk of tTGAs (HR: 1.30; 95% CI: 1.01 – 1.67), but tTGAs preceding IAs was not associated with risk of IAs (HR: 1.01; 95% CI: 0.69 – 1.50). Using a similar model adjusted for all other factors did not change this finding; IAs preceding tTGAs was significant (HR: 1.48; 95% CI: 1.15 – 1.91), whereas tTGAs preceding IAs was not (HR: 1.12; 95% CI: 0.75 – 1.67). The Cox Proportional Hazards model was then used to analyze demographic and genetic factors conferring risk of co-occurrence of IA and tTGA (Table 1). After adjustment for country, sex and having an FDR with CD, HLA-DR-DQ, HLA-DPB1 and the 5 SNPs at RG51, SH2B3, CTLA4, CCR3/CCR5, and PTPN2, we found HLA-DR3/4 (HR: 1.94; 95% CI: 1.12 – 3.35), having an FDR with T1D (HR: 2.69; 95% CI: 1.54 – 4.67) and rs3184504 at SH2B3 (HR = 1.53, 95% CI: 1.14 – 2.05) were each independently associated with co-occurrence of IAs and tTGAs.

**DISCUSSION**

The appearance of CD in existing T1D patients is widely appreciated. Our results analyze this relationship in a prospective manner at an earlier stage when autoantibodies to the respective diseases first develop. These autoantibodies are highly predictive of clinical disease and may be encountered more frequently as genetic screening and personalized medicine become more widespread. As with T1D and CD, observation in the first 5 years of life reveals that IAs precede tTGAs more often (67%) than vice versa (27%). Indeed, having IAs significantly increased the risk of developing tTGAs. The reverse relationship was not demonstrated, perhaps because of fewer individuals with the latter pattern detected during limited follow-up. Our results support the notion that children found to have IAs (for example through family screening or population-based prediction) should be screened for tTGAs, as is the practice for T1D children.

Having dual T1D and CD autoimmunity was more common than expected on the basis of the rates of each. It has been widely supposed that HLA antigen explains the overlap because similar
HLA-DR-DQ genotypes strongly predispose to both diseases. However, this did not appear to be the case. In a detailed analysis of the excess cases, considering HLA antigen loci, non-HLA antigen loci, and demographic features (FH, country) expected to capture additional (although unmeasured) genetic influences on these 2 autoimmune diseases, it is clear that not all the observed overlap between the 2 diseases was accounted for (Table 1). In addition to HLA-DR-DQ, SNP rs3184504 at SH2B3 accounted for significant overlap, but the effect was modest. SH2B3 is an adaptor protein involved in negative regulation of T-cells, consistent with both T1D and CD being T-cell–mediated diseases. Although not all common minor-effect variants or rare high-effect variants would be captured in the current analysis, our finding is in line with a previous study whose authors did not find major contributions by shared genetic risk factors for T1D and CD other than HLA antigen genotype. Our results therefore suggest that nongenetic factors are probably also involved.

Prospectively, IAs usually but not always preceded tTGAs, and previous IAs conferred a significantly increased risk of tTGAs. Although previous tTGA did not show a similar effect on subsequent IAs, there may have been insufficient individuals in this category for adequate evaluation. Our significant findings are in contrast to those of Williams et al, who found that tTGA prevalence in adults with an FH of T1D were not different between those with or without IAs, implying that tTGAs were not more likely in the setting of IAs. However, we prospectively managed individuals from early childhood when both types of autoimmunity first develop, which may explain our findings.

If preexisting T1D autoimmunity triggers CD autoimmunity, a possible mechanism might be coexpression of antigens from both diseases in the same anatomic location. tTGA is known to be expressed in islets. This stress leads to tTGA overexpression31 raises the intriguing possibility that tTGA expressed in the insulitis lesion, HAGOPIAN et al6

### Table 1: Cox Proportional Hazards Analysis of Factors Influencing Appearance of IAs and tTGAs, Their Order of Appearance, and Co-occurrence of IA and tTGA

<table>
<thead>
<tr>
<th>Model</th>
<th>Characteristics</th>
<th>IAs</th>
<th>tTGAs</th>
<th>Co-occurrence IAs and tTGAs</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>HR (95% CI)</td>
<td>P</td>
<td>HR (95% CI)</td>
</tr>
<tr>
<td>Unadjusted</td>
<td>tTGA preceding IA</td>
<td>1.01 (0.69–1.50)</td>
<td>.95</td>
<td>—</td>
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<tr>
<td></td>
<td>IA preceding tTGA</td>
<td>—</td>
<td>—</td>
<td>1.30 (1.01–1.67)</td>
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<tr>
<td>Adjusted</td>
<td>tTGA preceding IA</td>
<td>1.12 (0.75–1.67)</td>
<td>.57</td>
<td>—</td>
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<tr>
<td></td>
<td>IA preceding tTGA</td>
<td>—</td>
<td>—</td>
<td>1.48 (1.15–1.91)</td>
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<td>—</td>
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<tr>
<td></td>
<td>FIN</td>
<td>1.56 (1.21–2.00)</td>
<td>.0006*</td>
<td>1.24 (1.03–1.50)</td>
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<td></td>
<td>GER</td>
<td>1.14 (0.76–1.71)</td>
<td>.53</td>
<td>1.19 (0.97–1.48)</td>
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<td></td>
<td>SWE</td>
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<td>.023</td>
<td>1.57 (1.34–1.84)</td>
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<td></td>
<td>4/8</td>
<td>0.69 (0.50–0.96)</td>
<td>.028*</td>
<td>5.32 (4.32–6.56)</td>
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<td>DR 3/4</td>
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<td>2.05 (1.68–2.51)</td>
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<tr>
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<td>Female</td>
<td>0.76 (0.63–0.92)</td>
<td>.0039*</td>
<td>1.61 (1.41–1.84)</td>
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<tr>
<td>T1D FDR</td>
<td>No</td>
<td>2.60 (2.03–3.33)</td>
<td>&lt;.0001*</td>
<td>1.07 (0.84–1.34)</td>
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<td>Yes</td>
<td>1.01 (0.59–1.73)</td>
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<td>SH2B3 (rs156504)</td>
<td>1.30 (1.14–1.49)</td>
<td>&lt;.0001*</td>
<td>1.06 (0.97–1.17)</td>
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<td>PTPN2 (rs45450798)</td>
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<td>0.99 (0.87–1.12)</td>
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<td>.16</td>
<td>0.86 (0.74–1.02)</td>
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<td>HLA-DPB1*04:02</td>
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<td>0.93 (0.78–1.15)</td>
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<td>HLA-DPB1*01:01</td>
<td>1.03 (0.77–1.37)</td>
<td>.95</td>
<td>1.12 (0.94–1.34)</td>
</tr>
<tr>
<td></td>
<td>HLA-DPB1*02:01</td>
<td>1.21 (0.93–1.57)</td>
<td>.17</td>
<td>0.90 (0.73–1.10)</td>
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<td>HLA-DPB1*03:01</td>
<td>1.28 (0.98–1.68)</td>
<td>.08</td>
<td>1.08 (0.86–1.31)</td>
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</tbody>
</table>

Factors analyzed include country, sex, FH of T1D or CD, and HLA-DR-DQ, HLA-DPB1, and 5 non-HLA antigen genetic loci: FIN, Finland; GER, Germany; SWE, Sweden; US, United States; —, not applicable.

* P < .05
cotrafficking studies. Interestingly, a potential interaction between rotavirus was recently postulated to be reovirus was recently postulated to be with tTGA in TEDDY. Similarly, gluten exposure before 4 months increased CD risk in some but not all. It has been proposed that both T1D and CD are favored by an inflammation-mediated increase in gut permeability, which gluten is known to cause. However, age at introduction of gluten was not found to be significantly associated with tTGA in TEDDY. Gut microbes, in direct contact with the intestinal mucosa, are also prime candidates to influence CD risk. Rotavirus infection was implicated in both T1D autoimmune CD and CD autoimmunity, but both remain unconfirmed. These findings are further supported by TEDDY findings that a reported gastrointestinal infection 3 months before was significantly associated with tTGA seroconversion. Moreover, rotavirus vaccination had a protective effect in children exposed to gluten before 6 months of age, indicating a potential interaction between diet and infections on the risk of early autoimmunity. Interestingly, reovirus was recently postulated to be implicated in CD but its role in T1D is unknown. Gut bacterial microflora also differ between cases and controls for both CD and T1D, with cases of both diseases having fewer Gram-positive firmicutes and more Gram-negative bacteroides, although not all researchers agree and relationships are understandably complicated. The intestinal flora may therefore deviate in a similar way in both diseases. Future metagenomic analyses of TEDDY stool samples will test this possibility.

**CONCLUSIONS**

Early T1D and CD autoimmunity, their clinical diseases, occur together more than expected, with IAs usually preceding TTGAs. In addition to genes and demographics, nongenetic factors appear to contribute to co-occurrence. Elucidation of relevant mechanisms will require additional studies of local immune responses as well as common exposures (eg, gluten and microbial infections) within and outside of the TEDDY Study.

**THE TEDDY STUDY GROUP**

**Colorado Clinical Center**

Marian Rewers, MD, PhD, PI, Kimberly Bautista, Judith Baxter, Ruth Bedoy, Daniel Felipe-Morales, Kimberly Driscoll, PhD, Brigitte I. Frohnmert, MD, Patricia Gesualdo, Michelle Hoffman, Rachel Karban, Edwin Liu, MD, Jill Norris, PhD, Adela Samper-Imaz, Andrea Steck, MD, Kathleen Waugh, Hali Wright, University of Colorado, Anschutz Medical Campus, Barbara Davis Center for Childhood Diabetes.

**Finland Clinical Center**

Jorma Toppari, MD, PhD, PI, Olli G. Simell, MD, PhD, Annika Adamsson, PhD, Suvi Ahonen, Heikki Hyöty, PhD, Jorma Ilonen, MD, PhD, Sanna Jokipuu, Tiina Kallio, Leena Karlsson, Miia Kähönen, Mikael Knip, MD, PhD, Lea Kovanen, Mirva Koresalo, Kalle Kurppa, MD, PhD, Tiina Latva-aho, Maria Lönroth, Anette G. Ziegler, MD, PhD, Suvi Aho, Tiina Ninninen, Sari Niinrot, MD, PhD, youngster, Maria Vähä-Mäkelä, Mari Åkerlund, Katri Lindfors, PhD, University of Turku, University of Tampere, University of Oulu, Turku University Hospital, Hospital District of Southwest Finland, Tampere University Hospital, Oulu University Hospital, National Institute for Health and Welfare, Finland.

**Georgia/Florida Clinical Center**

Jin-Xiong She, PhD, PI, Desmond Schatz, MD, Diane Hopkins, Leigh Steed, Janey Adams, Katherine Silvis, Michael Haller, Melissa Gardiner, Richard McIndoe, PhD, Ashok Sharma, Joshua Williams, Gabriela Young, Stephen W. Anderson, Laura Jacobsen, Center for Biotechnology and Genomic Medicine, Augusta University.

**Germany Clinical Center:**

Anette G. Ziegler, MD, PhD, Andreas Beyerlein, PhD, Ezio Bonifacio PhD, Michael Hummel, MD, Sandra Hummel, PhD, Kristina Poterek, Nicole Janz, Mathilde Kersting, PhD, Annette Knopp, Sibylle Koletzko, Claudia Peplow, Roswith Roth, PhD, Marlon Scholz, Christiane Winkler, PhD, Forschergruppe Diabetes e.V. and Institute of Diabetes Research, Helmholtz Zentrum München, and Klinikum rechts der Isar, Technische Universität München. Center for Regenerative Therapies, TU Dresden, Dr von Hauner Children’s Hospital, Department of Gastroenterology, Ludwig Maximilians University Munich, Research Institute for Child Nutrition, Dortmund.

**Sweden Clinical Center**

Åke Lernmark, PhD, PI, Suvi M. Virtanen, MD, PhD, Mari Vähä-Mäkelä, Mari Åkerlund, Katri Lindfors, PhD, University of Turku, University of Tampere, University of Oulu, Turku University Hospital, Hospital District of Southwest Finland, Tampere University Hospital, Oulu University Hospital, National Institute for Health and Welfare, Finland.
Co-occurrence of Type 1 Diabetes and Celiac Disease

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Andrén Aronsson,2,12,13 Maria Ask, Jenny Bremer, Ulla-Mari Carlsson, Corrado Cilio, MD, PhD,6 Emelie Ericson-Hallström, Lina Fransson, Thomas Gard, Joanna Gerardsson, Rasmus Bennet, Monica Hansen, Gertie Hansson, Susanne Hyberg, Fredrik Johansen, Berglind Jonsdottir, MD, Helena Elding Larsson, MD, PhD,6,14 Marielle Lindström, Markus Lundgren, MD,14 Maria Månsson-Martínez, Maria Markan, Jessica Melin,12 Zeliha Mestan, Karin Ottosson, Kobra Rahmati, Anita Ramelius, Falastin Salami, Sara Sibthorpe, Birgitta Sjoberg, Ulrica Swartling, PhD,9,12 Evelyn Tekum Amboh, Carina Törn, PhD,3,15 Anne Wallin, Åsa Wimar,12,14 Sofie Åberg, Lund University.

Washington Clinical Center


Pennsylvania Satellite Center

Dorothy Becker, MD, Margaret Franciscus, MaryEllen Dalmagro-Elias Smith,2 Ashi Daftyari, MD, Mary Beth Klein, Chrystal Yates. Children’s Hospital of Pittsburgh of UPMC.

Data Coordinating Center

Jeffrey P. Krischer, PhD, PI,1,4,5,10,11 Michael Abbondondolo, Sarah Austin-Gonzalez, Maryouri Avendano, Sandra Baethke, Rasheedah Brown,12,15 Brant Burkhart, PhD,5,6 Martha Butterworth,2 Joanna Clasen, David Cuthbertson, Christopher Eberhard, Steven Fiske,9 Dena Garcia, Jennifer Garmeson, Veena Gowda, Kathleen Heyman, Francisco Perez Laras, Hye-Seung Lee, PhD,12,13,15 Shu Liu, Xiang Liu, PhD,2,3,9,14 Kristian Lynch, PhD,5,6,9,15 Jamie Malloy, Cristina McCarthy,12,15 Steven Meulemans, Hemang Parikh, PhD,2,3 Chris Shaffer, Laura Smith, PhD,9,12 Susan Smith,12,15 Noah Sulman, PhD, Roy Tamura, PhD,12,13 Ulla Uusitalo, PhD,2,15 Kendra Vehik, PhD,4,6,14,15 Ponni Vijayakandipan, Keith Wood, Jimin Yang, PhD, RD,2,15 Past staff: Lori Ballard, David Hadley, PhD, Wendy McLeod. University of South Florida.

Project Scientist

Beena Akolkar, PhD,1,3–7,10,11 National Institutes of Diabetes and Digestive and Kidney Diseases.

Other Contributors

Kasia Bourcier, PhD,5 National Institutes of Allergy and Infectious Diseases. Thomas Briese, PhD,6,15

Columbia University. Suzanne Bennett Johnson, PhD,9,12 Florida State University. Eric Triplett, PhD,6 University of Florida.

Committees

1Ancillary Studies, 2Diet, 3Genetics, 4Human Subjects/Publicity/Publications, 5Immune Markers, 6Infectious Agents, 7Laboratory Implementation, 8Maternal Studies, 9Psychosocial, 10Quality Assurance, 11Steering, 12Study Coordinators, 13Celiac Disease, 14Clinical Implementation, 15Quality Assurance Subcommittee on Data Quality.

ABBREVIATIONS

CD: celiac disease
CI: confidence interval
FD: first-degree relative
FH: family history
HR: hazard ratio
IA: islet autoantibodies
IAA: insulin autoantibodies
IQR: interquartile range
SNP: single-nucleotide polymorphism
T1D: type 1 diabetes
TEDDY: The Environmental Determinants of Diabetes in the Young
TtTG: tissue transglutaminase
TtGA: tissue transglutaminase autoantibodies

and supervise the single-nucleotide polymorphism genotyping effort, secured the funding, and reviewed and revised the manuscript; Dr Krischer helped conceptualize and design the study, secured funding, coordinated the study and its central data storage, and reviewed and revised the manuscript; Dr Erlich helped conceptualize, design, and conduct the HLA antigen genotyping effort and reviewed and revised the manuscript; Dr Akolkar helped conceptualize, design, and fund the study, coordinated the genetic typing parts of the study, and reviewed and revised the manuscript; each author has participated sufficiently in the work to take public responsibility for appropriate portions of the content; and all authors approved the final manuscript as submitted and agree to be accountable for all aspects of the work.

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Address correspondence to William Hagopian, MD, PhD, Pacific Northwest Research Institute, 720 Broadway, Seattle, WA 98122. E-mail: wah@uw.edu

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REFERENCES


