1 TITLE

2	Models for infantile hypertrophic pyloric stenosis development in patients with esophageal atresia
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30	Summary statement:
31 32	Instead of one affected gene, the higher incidence of IHPS in EA patients is more likely the result of multiple (epi)genetic and environmental factors together shifting the balance to disease

33 development.

34 ABSTRACT

35 Patients born with esophageal atresia (EA) have a 30 times higher prevalence of infantile 36 hypertrophic pyloric stenosis (IHPS). This makes sense from a developmental perspective as both the 37 esophagus and the pyloric sphincter are foregut derived structures. EA and IHPS are variable features 38 in several (monogenetic) syndromes. This, and twin and familial studies, indicates a genetic 39 component for both conditions as single entities. We hypothesized that genetic defects, disturbing 40 foregut morphogenesis, are responsible for this combination of malformations. Non-genetic factors 41 could also contribute, as mice exposed to Adriamycin develop EA and in utero diethylstilbestrol 42 exposure is associated with EA.

We investigated the copy number profiles and protein coding variants of 15 patients with both EA and IHPS. As all parents were unaffected, we first considered dominant (*de novo*) or recessive inheritance models but could not identify putatively deleterious mutations or recessive variants. We did identify inherited variants in genes either known to be involved in EA or IHPS or important in foregut morphogenesis in all patients. Unfortunately, variant burden analysis did not show a significant difference with unaffected controls. However, the IHPS associated risk SNP rs1933683 had a significantly higher incidence (OR 3.29, p=0.009).

Although the genetic variation in likely candidate genes as well as the predisposing locus near *BARX1* (rs1933683) suggest a genetic component, it does not fully explain the abnormalities seen in these patients. Therefore, we hypothesize that a combination of high impact genetic, mechanical and environmental factors together can shift the balance to abnormal development.

54 INTRODUCTION

55 Esophageal atresia (EA) is a rare congenital malformation caused by a faulty development of the 56 foregut which leads to a discontinuity of the esophagus. It occurs in about 2.5 cases per 10,000 births 57 within Europe (Pedersen et al., 2012, Oddsberg et al., 2012) and over three-quarters of patients 58 present with a tracheoesophageal fistula (TEF) (Pedersen et al., 2012, Macchini et al., 2017). EA is 59 considered - etiologically as well as phenotypically - a highly heterogeneous condition (Brosens et 60 al., 2014). It can present either as an isolated defect but is often seen in combination with other 61 malformations. Frequently, these malformations are part of the VACTERL (Vertebral, Anorectal, 62 Cardiac, Tracheoesophageal, Renal or urinary tract of Limb malformations) association. VACTERL 63 association is a diagnosis of exclusion in which three or more features of the VACTERL spectrum are 64 present and a known genetic syndrome is not identified (Solomon et al., 2012). However, clustering 65 of one or more of these features with additional specific associated malformations could also be the 66 results of a shared genetic etiology.

One of the more prevalent, but less well-known, associated malformations is Infantile Hypertrophic Pyloric Stenosis (IHPS) (Rollins et al., 1989). In contrast to EA, IHPS is often considered an acquired disorder. The pyloric muscle hypertrophies in the first weeks of life, causing a narrowing of the pyloric channel (Panteli, 2009). Healthy-born infants present at week 3 to 6 of life with projectile postprandial vomiting. They need surgery where the upper layer of the circular smooth muscle of the pylorus will be incised, to release the passage from the stomach to the intestine again.

73 Previously, we described a 30 times higher prevalence (7.5%) of IHPS in patients with EA 74 compared to the normal population (0.25%) (van Beelen et al., 2014). This increased prevalence has 75 been reported in other retrospective studies (3.3-13%) as well (Palacios M.E.C. et al., 2014, Deurloo 76 et al., 2002). The clinical presentation of IHPS seen in patients with EA/IHPS is not different from 77 patients with isolated IHPS. However, the diagnosis of IHPS is more difficult and often delayed in 78 patients with EA. Relatively common complications after EA repair, such as stenosis of the 79 anastomosis, can protect against reflux and lead to just regurgitation. By the time these patients 80 start vomiting, there must be massive gastroesophageal reflux.

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The presentation of both EA and IHPS makes sense from a developmental perspective as the esophagus and the pyloric sphincter are both foregut derived structures. Organ specification during embryonic development is under tight spatiotemporal control of specific growth factors, transcription factors and signaling cascades (Li et al., 2009, Jacobs and Que, 2013). Disturbances in these pathways could impact proper development. In mice, the esophagus is specified from the foregut tube between embryonic day E9.5 and E11.5. In humans, the esophagus, as well as the

stomach, starts developing from the fourth week after conception onwards. The stomach turns around its anterior-posterior axis during embryonic development (Cetin et al., 2006). The developing pylorus can be visualized with immunostaining at week six after gestation and differentiates during fetal life (Koyuncu et al., 2009).

92 Environmental (Zwink et al., 2016, Felix et al., 2008, Feng et al., 2016, Markel et al., 2015, Krogh et 93 al., 2012, Sorensen et al., 2002) and genetic contributions (Peeters et al., 2012, Brosens et al., 2014, 94 Solomon et al., 2012) have been described for both EA and IHPS as single entities or in combination 95 with other anatomical malformations. It has been suggested that in utero exposure to 96 diethylstilbestrol (DES) is associated with the development of EA (Felix et al., 2007a). Moreover, both 97 malformations are variable features in specific and often phenotypically overlapping genetic 98 syndromes (Table 1). The presence of both conditions as variable features in the phenotypical 99 spectrum of known genetic syndromes is indicative of a genetic background for EA and IHPS. More 100 evidence for a genetic contribution can be deduced from twin studies and animal models (de Jong et 101 al., 2010). The concordance rates in monozygotic twins compared to dizygotic twins is higher for EA 102 (Veenma et al., 2012) and IHPS (Krogh et al., 2010) as single entities. Also, the recurrence risk is 103 elevated for siblings and offspring of affected individuals with EA in combination with other 104 associated anomalies (Robert et al., 1993, Van Staey et al., 1984, Warren et al., 1979, McMullen et 105 al., 1996). In contrast, the recurrence risk for isolated EA is low (Schulz et al., 2012) and moderate for 106 IHPS (Krogh et al., 2010, Elinoff et al., 2005). In contrast to EA, there has been reported a male 107 predominance for IHPS (4:1) (MacMahon, 2006). There have been risk loci associated to IHPS 108 (Everett and Chung, 2013, Feenstra et al., 2012, Feenstra et al., 2013, Svenningsson et al., 2012, 109 Fadista et al., 2019). To date, no risk loci have been described for EA.

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111 Considering the increased prevalence of IHPS in patients with EA, their common developmental 112 origin and previous evidence in genetic studies, we hypothesized that Copy Number Variants (CNVs) 113 or other protein coding alterations affecting one specific gene, or genetic disturbances in more genes 114 all important for foregut morphogenesis are responsible for the higher incidence of IHPS in patients 115 with EA.

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- 117

118 **RESULTS**

119 Patient cohort

120 In total, 27 out of 664 patients (4.1%) born with EA between 1970-2017, developed IHPS. Twenty 121 patients have been described previously (van Beelen et al., 2014). Parental informed consent for

whole exome sequencing (WES) was obtained for 15 patients. Several phenotypical characteristics stood out in this EA/IHPS cohort: a sacral dimple was present in seven patients (25.9%), anomalies of the vertebrae or ribs in eight patients (29.7%) and genitourinary anomalies in six patients (22.2%) of which two patients (7.4%) had hypospadias. Four patients (14.8%) had three or more anomalies within the VACTERL spectrum (Solomon, 2011). A full phenotypical description of the 27 EA/IHPS patients is given in Table 2.

128

129 Copy Number analysis

130 Our previous study described rare CNVs and their inheritance pattern in patients with EA (Brosens et 131 al., 2016b), seventeen EA/IHPS patients were included in this previous study. None of the six large 132 CNVs identified were *de novo*, all were inherited from one of the unaffected parents. Patient 133 SKZ_400 had a paternal inherited rare gain of chromosomal region 11q15. Patient SKZ_0887 had 134 maternal inherited putative deleterious gains on Xq26.1 and Xp22.33. Patient SKZ 1003 had a 135 maternal inherited loss of chromosomal region 17q11 and patient SKZ 1248 maternal inherited rare 136 gains in chromosomal regions 4q35 and 5p15.1. Additional exon-level CN-profiling using the 137 normalized coverage profiles (Amarasinghe et al., 2013) of the exome sequencing data confirmed the 138 presence of the CNV seen with SNP-array. All CN profiles of main EA and IHPS disease genes (Brosens 139 et al., 2014, Peeters et al., 2012) were normal. There were no overlapping rare CNVs in this patient 140 cohort. All rare CNVs, classified as VUS or (likely) deleterious are described in Table S1.

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142 Exome sequence analysis

Sequencing resulted in at least 5 Giga-bases of raw sequence data with an average coverage of 70X and 90% of target bases covered over 20X. Quality of the sequence data is listed in Table S2. As none of the parents of the 15 investigated patients were affected we first considered dominant *de novo* and recessive modes of inheritance.

147 We could not identify de novo pathogenic variation in main EA and IHPS disease genes (Brosens et 148 al., 2014, Peeters et al., 2012). Subsequently, we searched for possible de novo mutations exome 149 wide. For this, we focused on putative deleterious ultra-rare protein coding or splice site variants 150 (n=100) (Bennett et al., 2017). Variations were considered ultra-rare when they were absent in the 151 gnomAD dataset (123,136 whole exomes and 15,496 whole genomes) 152 (<u>http://gnomad.broadinstitute.org/</u>) (Lek et al., 2016). Twenty-five variants proved to be sequencing 153 artifacts. Furthermore, we could not confirm the segregation of 15 mutations due to lack of parental 154 DNA. We determined the segregation of all ultra-rare variants predicted to be of unknown 155 significance (VUS, n=37) or (likely) deleterious (n=23). All putative deleterious variants tested proved 156 to be inherited from one of the unaffected parents.

157 Considering a recessive mode of inheritance, we searched for genes with homozygous or 158 compound heterozygous variants. Six variants in three genes (FLNC, ATP6V0A1 and FAM46A) fitted a 159 putative compound heterozygous model, two genes (KCNN3 and VDAC3) had homozygous variants 160 and two genes (MID2 and SH3KBP1) had variants on chromosome X in a male patient. All variants 161 were predicted to be likely deleterious or VUS and intolerant to missense variants (Z-score \geq 3) or loss 162 of function variants (PLI or PLIrec \geq 0.9). With segregation analysis, we could confirm the compound 163 heterozygous mode of inheritance of the variant in the FAM46A gene in patient SKZ 2023 and the 164 maternally inherited X-linked variant in the SK3KBP1 gene in patient SKZ 1260. The other recessive 165 candidate genes could not be validated due to technical difficulties (and are likely sequencing 166 artifacts) or due to lack of parental DNA. None of the recessive candidate genes were affected twice 167 or more in this cohort. All predicted deleterious variants were submitted to the ClinVar database 168 https://www.ncbi.nlm.nih.gov/clinvar/ (Landrum et al., 2014).

We inspected the CN profiles from WES-CN and SNP-array for partial overlap with genes affected by heterozygous variant predicted to be deleterious in (recessive) loss of function intolerant or missense intolerant genes (n=48) and could not detect unmasking of a recessive mutation by a CNV. Ultra-rare variants (n=78), X-linked or recessive variants are depicted in Table S4 and uploaded to the ClinVar database (https://www.ncbi.nlm.nih.gov/clinvar/)

- 175 Chirvar uatabase (<u>https://www.httpi.him.him</u>
- 174

175 Pathway enrichment analysis of genes affected by rare variants

176 When looking at the selected protein altering variants (Z-score \geq 3, n=44) or loss of function 177 intolerant (PLI \geq 0.9, n=4), two relevant pathways were significantly enriched (p-value <1x10⁻⁵): 178 proliferation and differentiation of smooth muscle cells (*INSR*, *ITGB1*, *NOTCH1*, *TCF4*, *PDE4D*, *TERT*, 179 *ANKRD17*, *DICER1*) and self-renewal of satellite cells (*ITGB1*, *NOTCH1*).

180

181 Variant prioritization using different in silico tools

We prioritized all rare variants with three in silico tools (see Methods section). Fifty-four variants in
34 genes had an overlap between VAAST (Yandell et al., 2011, Hu et al., 2013, Kennedy et al., 2014)),
which prioritizes based on variant deleteriousness and Phevor and PhenIX which prioritize more on
phenotype (Singleton et al., 2014, Zemojtel et al., 2014)). Top ranking variants can be found in Table
S3.

Additionally, we found variants in the same gene in multiple patients (Fig. 1). Of these 116 genes (VUS=87, likely deleterious=30), 36 genes were found in \geq 3 patients of which six genes were present in more than five patients (*CNTN2*, *DSPP*, *NOTCH4*, *PRRC2A*, *SEC16B*, *ZNF717*). Four (*AMBRA1*, *ATP2A3*, *DSCAM*, *NOTCH1*) out of 116 genes were predicted to be intolerant for missense variants (Zscore \geq 3). See also Table S4.

192

193 Gene burden analysis

194 An exome wide gene burden analysis showed ten genes which were enriched for rare putatively 195 deleterious variation compared to the 1000 Genomes project phase 3 samples. The results are 196 shown in Table 3. There were no genes with more than two distinct variants. Each variant was 197 observed only once. A second burden test - only evaluating genes from developmental important 198 pathways and known disease genes – showed no significant difference between our 15 patients and 199 a control group of 44 healthy individuals, who were sequenced in a previous study (Table 4). Also, 200 the number of putative deleterious variants between these two groups was not significantly different 201 (Table 5). Unfortunately, a burden test comparing the variant profiles of these genes between the 202 patients and their parents was not possible since no WES data of the parents was available.

203 Expression of main candidate gene during development

With public micro-array transcriptome data we evaluated which genes were upregulated at a specific
time-point in the foregut, esophagus or pyloric sphincter and used the output as an indicator of gene
expression (see Methods section and Table S5). Of the genes classified as VUS or likely deleterious in
our exome sequencing results, 28 genes were upregulated in both the foregut or esophagus as well
as the pyloric sphincter: *ADAMTSL4, AGRN, ANKRD29, ARHGAP29, CAMTA1, CDHR5, CNTN2, COL11A1, DNAJC11, HIVEP3, HMCN1, HMGCS2, HSPG2, ITGB3BP, LDB3, MYOF, NKX2-3, NUP133, PCSK9, PKN2, PRDM16, PUM1, RET, SEC16B, SERINC2, TMEM82, VPS13D* and *ZBTB7B*.

Unfortunately, none of the genes enriched in our burden analysis were differentially expressed in mice foregut between E8.5 and E16.5. Seven out of 116 genes with putative deleterious variants in more than one patient were differentially expressed in mice foregut: *Adamtsl4* at E8.5, E14.5 and E16.5; *Ankrd26* at E14.5; *Cntn2* at E8.5, E15.5 and E18.5; *Hspg2* at E8.25, E8.5, E14.5 and E18.5; *Kcnn3* at E8.5 and E15.5; *Ldb3* at E8.5, E14.5 and E15.5; *Sec16b* at E8.5, E14.5 and E16.5. Of the top candidate genes in the manual burden analysis (see Table 4 and Table S6) only *Ret* was differentially expressed in mice at E8.25, E8.5, E11.5, E14.5, E15.5, E16.5 and E18.5.

218

219 Detection of common SNPs associated with IHPS

Determination of the risk allele frequency of four loci highly associated with IHPS (rs11712066, rs573872, rs29784 and rs1933683 near genes *MBNL1*, *NKX2-5* and *BARX1*, respectively) revealed a significantly higher incidence of rs1933683 in our EA/IHPS cohort compared to the population frequency (OR 3.29 (95% CI 1.27-8.56), p=0.009, see Table S7). The risk allele frequency of the other risk loci was not significantly different from the normal population. We did not detect rare putatively deleterious variants in *MBNL1*, *NKX2-5* and *BARX1* in the patient exome sequencing data.

226 **DISCUSSION**

We hypothesized that the increased prevalence of IHPS in patients with EA compared to the prevalence of IHPS in the normal population was due to shared CNVs or protein coding alterations in a specific gene, or due to genetic disturbances in genes of shared biological networks during development. As mentioned earlier, both EA and IHPS are variable features in specific genetic syndromes (Table 1). Therefore, to find genetic aberrations that contribute to EA/IHPS we initially searched for pathogenic alterations in known EA or IHPS associated genes (Table S6).

233

234 There are no pathogenic changes in known disease genes

As all parents were unaffected, we started this study by focusing on *de novo*, recessive or X-linked changes affecting these known disease genes. However, we could not identify deleterious protein coding alterations, exonic gains or losses or larger CNVs affecting these genes. This is in line with previous studies in which limited causal changes could be detected in patients with EA and associated anomalies (Zhang et al., 2017, Hilger et al., 2015, Brosens et al., 2016b)

240 Next, we extended our analysis to all genes covered in the exome capture. Given the small sample 241 size (n=15), the low prevalence of the disorder and the high impact on development, we 242 concentrated on genes intolerant to variation (Lek et al., 2016, Ruderfer et al., 2016) harboring rare 243 putative deleterious single nucleotide changes or large CNVs. Moreover, we determined the 244 segregation of alterations in these candidate genes, putative recessive (X-linked, compound 245 heterozygous and homozygous recessive) and all ultra-rare protein altering changes absent from the 246 gnomAD database (Lek et al., 2016) as the later have a high chance of being de novo (Bennett et al., 247 2017). Unfortunately, we did not identify any de novo mutations or de novo CNVs. None of the 248 identified inherited rare CNVs overlapped in these patients. We could confirm the presence of a 249 compound heterozygous variant in FAM46A in one patient and an X-linked variant in SH3KBP1 in 250 another patient. However, FAM46A and SH3KBP1 are not known to be associated with the 251 gastrointestinal or respiratory tract and were not differentially expressed at the time points 252 important for foregut morphogenesis. These findings made us conclude that neither a dominant nor 253 a recessive model can explain the combination of EA and IHPS in these patients.

254

255 The coding sequences of genes crucial in esophageal and pyloric sphincter formation are affected

Subsequently, we focused on genes involved in foregut development. Literature research together
with data of previous expression studies provided an overview of genes important for foregut
development (Fig. 2).

259 The development of the foregut is most studied in mouse models. In mice, early foregut 260 formation starts with Foxa2 stimulation of the anterior endoderm at E8.0 (Heath, 2010). The 261 endodermal sheet folds and forms a tube at E8.75 (Sherwood et al., 2009). Next, signals from the 262 notochord start dorsal-ventral patterning around E9.0, with high Nkx2.1/absent Sox2 in the ventral 263 future trachea and absent Nkx2.1/high Sox2 in the dorsal future esophagus and stomach (Que et al., 264 2007). These dorsal-ventral patterns lead to compartmentalization of the foregut. Between E9.5 and 265 E11.5 the foregut separates in the primordial esophagus and stomach, and in the primordial trachea. 266 Primordial lung buds become apparent at E9.5 (Sherwood et al., 2009). The separation site is marked 267 by mesenchymal expression of Barx1 (Woo et al., 2011). The esophagus is completely separated from 268 the trachea at E11.5.

269 Pyloric sphincter formation is mostly studied in chick and mouse models. This formation starts 270 with the thickening of the circular smooth muscle layer between the antrum and the duodenum 271 around E14.5 and the primordial pyloric sphincter is complete around E18.5 (Smith et al., 2000a, Self 272 et al., 2009). In addition to its functioning in foregut separation, the *Barx1* homeobox gene is also 273 vital for stomach differentiation and stomach smooth muscle development. It inhibits Wnt signaling 274 (Woo et al., 2011) and modulates the expression of Bapx1, another important factor required for 275 pyloric sphincter morphogenesis (Jayewickreme and Shivdasani, 2015, Stringer et al., 2008, Verzi et 276 al., 2009).

277

278 Given the described importance of these genes in normal development, we hypothesized that 279 variations in multiple genes important for foregut morphogenesis might explain the higher incidence 280 of IHPS in patients with EA. We compared a selection of genes – known to be important for foregut 281 morphogenesis or syndromatically associated with EA or IHPS – between the patients and the 282 healthy controls (Table 4, Table S6). Interestingly, in TNXB (NM 019105.6:c.4444G>A, 283 p.Val1482Met), WDR11 (NM 018117.11:c.1138G>T, p.Val380Phe), PEX3 (NM 003630.2:c.1012A>G, 284 p.Ser338Gly), TBX3 (NM_016569.3:c.506G>A, p.Arg169Gln), and GDF6 (NM_001001557.2:c.281C>G, 285 p.Pro94Arg) rare variants were present that were absent in the control group. These variants might 286 not be sufficient to result in disease but are predicted to impact the protein and might contribute 287 together with other unknown factors to disease development.

288

289 There is no high frequency burden of rare variants

Given the limited number of samples, we will only detect a gene burden if it is large and has a high impact. We compared the total rare and ultra-rare variant burden of putative deleterious variants in all genes. The number of ultra-rare variants was slightly higher in the patient group compared to the control group but did not differ significantly (Table 5). A second burden analysis identified ten genes

with more variants compared to those seen in the 1000 Genome cohort, two variants were predicted to be deleterious (Table 3). Unfortunately, they did not show any overlap with the results of the expression analysis or candidate genes selected from the literature. Therefore, these variants are not likely to explain the increased incidence of IHPS in EA patients or EA/IHPS development. A rare variant burden might exist but we could not detect it due to limited sample size and/or focus on known candidate genes.

300

301 Of all the protein coding changes classified as VUS or higher (Table S4; Table S8), 116 genes were 302 affected with a variant in more than one patient (Fig. 1). Seven of these genes (ADAMTSL4, 303 ANKRD26, CNTN2, HSPG2, KCNN3, LDB3, SEC16B) were differentially expressed in the developing 304 foregut, esophagus or pyloric sphincter in mice between E8.25 and E16.5. However, none of these 305 genes could explain the combination of EA and IHPS within a patient based on their function; none of 306 these genes is known to be associated with the gastrointestinal or respiratory tract. Furthermore, 307 most variants had a population frequency above the prevalence of EA/TEF. If these variants are 308 highly penetrant, they would not be the likely cause. Increasing sample sizes (drastically) would allow 309 an analysis going beyond known intolerant genes, allow us to consider reduced penetrance and 310 potentially identify a shared genetic etiology.

311

312 Known common variants associated with IHPS development could have an impact in some patients

313 Since certain SNPs have been identified with GWAS to be highly associated with IHPS, we wondered 314 if these known common haplotypes could also play a role in the higher incidence of IHPS in patients 315 with EA. In our cohort, we found a significantly higher incidence of the risk loci rs1933683 compared 316 to the population frequency (Table S7). Three patients were homozygous for the risk allele and have 317 a substantially increased risk for IHPS development. The common risk haplotype might therefore 318 impact IHPS development in some of the IHPS patients. However, further research is needed to 319 confirm the impact of this haplotype in a larger EA and EA/IHPS population.

320 Possible contribution of non-genetic factors

All the data presented so far made us conclude that dominant *de novo* variations in possible disease causing genes do not play a role in our cohort. Recessive inheritance cannot totally be excluded, although our results are not suggestive for this mode of inheritance. We did identify in all patients putative disease-causing variants. Nevertheless, as all parents from whom these variants were inherited were not affected, these variants could contribute but not cause the disease. Previous studies suggested the contribution of non-genetic factors as an explanation for the combined occurrence of EA and IHPS.

328 Could IHPS be an acquired condition related to surgery or treatment of EA?

329 The overrepresentation of IHPS in EA patients made us wonder if IHSP could also be the result of the 330 atresia itself, potentially as a result of the surgical procedure to correct the atresia or the result of 331 treatment. Previous studies also mentioned vagal nerve lesions, a gastrostomy and transpyloric 332 feeding tubes as possible causes for an increased incidence of IHPS after correction of EA (Ilhan et al., 333 2018). IHPS has been suggested to be a neuromuscular disorder with the involvement of smooth 334 muscle cells, interstitial cells of Cajal and the enteric nervous system. The hypertrophy is suggested 335 to be the result of discoordinated movements of the pyloric sphincter and the contractions of the 336 stomach (Hayes and Goldenberg, 1957), perhaps as the result of absent nitric oxide synthase activity 337 (Vanderwinden et al., 1992). Impaired gastric contractility and esophageal relaxation were observed 338 in Adriamycin and doxorubicin induced EA in mice (Tugay et al., 2003, Tugay et al., 2001). 339 Mechanistically, this association between EA and IHPS seems plausible. However, it does not explain 340 why IHPS is not fully penetrant in patients with EA. The most common thought is that mechanical 341 and environmental factors disturb the developmental field. To which extent these factors influence 342 the development of the child, depends on the specific risk factors and their timing. Further research 343 on the cause and other specific clinical risk factors for patients with EA should be considered, e.g. the 344 late start of oral feeding or the long-term feeding through a tube instead of drinking themselves.

345

346 Models for EA/IHPS disease etiology

Since we hypothesized that genetic defects, disturbing foregut morphogenesis, would be responsible for the combination of EA/IHPS, we started with the thought of a (monogenetic) syndromic model. However, we have not been able to find a central gene impacted in most patients which can explain the increased prevalence of IHPS in patients with EA. An as yet unknown syndrome is unlikely since we have not found *de novo* (as all parents are unaffected) or shared high impact variants in the same gene multiple patients. However, we cannot exclude *de novo* mutations which have been seen in the GnomAD exomes or genomes, nor did we look beyond the coding part of our genome.

354 Furthermore, we have detected inherited rare variants in candidate genes and genes affected 355 more than once by variants with a low (unaffected) population frequency. Therefore, we cannot 356 exclude a genetic component. Another option is that, although the combination of EA and IHPS could 357 not be explained by one gene or locus, the underlying cause of EA and the other associated 358 anomalies is the result of hits in multiple genes. About 10% of the patients with EA have an 359 underlying genetic syndrome (Brosens et al., 2014) and more can be expected. The phenotypical 360 spectrum of this cohort is very heterogeneous and could be the result of impacts on multiple genes. 361 IHPS could then independently be caused by mechanical factors such as the surgical procedure.

362

363 Furthermore, environmental risk factors have been suggested for EA and IHPS, like pesticides, 364 smoking, herbicides and periconceptional alcohol or multivitamin use (Zwink et al., 2016, Felix et al., 365 2008, Feng et al., 2016, Markel et al., 2015, Krogh et al., 2012, Sorensen et al., 2002). Considering the 366 absence of highly penetrant recurring genetic variations, we now hypothesize different multifactorial 367 models for disease development. In all these models the combination of CNVs, deleterious protein 368 alterations (Felix et al., 2007b, Brosens et al., 2014), severe changes in the developmental field 369 during the organogenesis (Martinez-Frias, 1994, Martinez-Frias and Frias, 1997) and/or 370 environmental inducing epigenetic changes (Sorensen et al., 2002) together can modulate the 371 phenotypical spectrum seen in these patients. Examples of possible mechanical and environmental 372 factors disturbing the developmental field are mice models with Adriamycin induction or dorsal-373 ventral patterning signals from the notochord.

374

375 Our first hypothesis is based on the earlier published theory of Brosens et al. about disturbed ENS 376 development (Brosens et al., 2016a). It includes a seesaw model in which risk factors are in balance 377 with protective mechanisms. In this model, the fulcrum can be shifted by a genetic variation in a 378 central gene, which automatically disrupts the balance. When applying this theory on EA/IHPS 379 patients, this would lead to more or less affected organ systems within the VACTERL syndrome. 380 However, in this study we have not been able to detect one central gene. This makes a seesaw model 381 with a variable fulcrum less likely as an explanation for the increased prevalence of IHPS in the EA 382 population.

383

384 A second hypothesis is a burden model (Fig. 3A). Similar, (epi)genetic, environmental and mechanical 385 factors form a burden of risk factors, which balances with protective mechanisms. In this model, the 386 point of balance is not shifted by a mutation in a central gene and every person has contributions of 387 certain risk factors. But in most cases this does not lead to affected organ systems. There is an 388 intermediate range between normal and affected in which individuals can have the genetic burden 389 but lacks an abnormal phenotype (reduced penetrance) or their symptoms differ in severity (variable 390 expressivity). The latter would fit the results in this study; maybe we did have detected variants but 391 have we failed to interpret them correctly as parents were seemingly unaffected and/or the variant 392 frequency can be higher in unaffected controls to be of relevance in patients. Mechanical or 393 environmental factors could have made the difference in shifting the balance. All together the 394 burden model is a plausible explanation for the disease development.

395

Last, we hypothesize a slippery slope model (Fig. 3B). In this model, the burden of low impact geneticvariants and environmental disturbances alone does not impact the balance seen in the seesaw

398 model unless it crosses a certain threshold. Moreover, we hypothesize that the protective 399 mechanisms (e.g. compensatory mechanisms) during development are very strong, making it really 400 difficult to shift the balance. Most fetuses do not develop any malformations despite the combined 401 genetic and environmental burden or do not survive. But once the threshold is reached, the balance 402 is immediately greatly disrupted and often multiple organ systems are affected. This model fits with 403 the phenotypical results in this study since four patients (14.8%) had three or more anomalies within 404 the VACTERL spectrum. In this model there is a high tolerance for low impact genetic variation and 405 only high impact variation (aneuploidies, exposure to toxic substances, pathogenic changes in 406 developmental crucial genes) shifts the balance. When the balance is disturbed, it shifts drastically. 407 We did not detect high impact changes responsible for the EA/IHPS combination. As parents are 408 unaffected it is (in this model) unlikely that inherited variants impact disease development, nor 409 would variants which are seen in the (unaffected) population controls.

410

411 Limitations

412 Not finding any positive correlation between DNA variations in specific genes or developmental 413 pathways is partly due to the small data set we have (15 patients). The small sample size is no 414 problem for the *de novo* and our recessive model strategy in known disease genes, but it is so for the 415 heterozygous variant burden analysis. Another limitation is the lack of data on the expression of 416 genes involved in normal foregut development in human embryos. Our gene selection was based on 417 mouse transcriptome data. Little human data is available since human embryos of 4 to 6 weeks old 418 are generally not preserved. However, although it is unclear how precisely the foregut development 419 in mice corresponds with humans, it is unlikely that this is very different in its early phases. Finally, 420 one could argue that variations in the non-coding part of the genome are major contributors. 421 Although we did investigate known IHPS risk loci and determined genome wide CNV profiles, we did 422 not determine genome wide variation.

423

424 Conclusions

To conclude, *de novo* mutations (a dominant model) and homozygous or compound heterozygous mutations (a recessive model) in the protein coding part of the genome are not a likely cause for the combination of EA and IHPS. Although the presence of genetic variation in likely candidate genes suggests a genetic component, there does not seem to be an enrichment of genetic variants in good candidate genes in patients. There are putative deleterious variants in foregut or disease genes which might contribute to disease development and although there is no difference in burden, some variants might contribute more than others and this is not taken into account in a burden test. We 432 might have misinterpreted the impact of some of the inherited variants. Furthermore, in some433 patients the IHPS predisposing locus rs1933683 is present.

We hypothesized several multifactorial models in which the combination of multiple high impact genetic, mechanical and environmental factors together can shift the balance from normal to abnormal development. A burden model with reduced penetrance or variable expressivity is most likely if genetic factors contribute. Future research should investigate the incidence of IHPS in bigger EA patients cohorts to further explore this theory. To exclude the role of treatment or surgery, clinical factors related to the surgical correction of EA – for example vagal nerve lesions after surgery, the late start of oral feeding or transpyloric feeding tubes – should be systematically registered.

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- 442

443 MATERIAL AND METHODS

444 Patient cohort

This study was approved by the Medical Ethical Review Board of Erasmus MC - Sophia Children's Hospital (MEC 193.948/2000/159). We searched the Erasmus University MC-Sophia EA-cohort and the database of the standardized prospective longitudinal follow up program in our hospital for children with congenital anatomical anomalies (Gischler et al., 2009) for patients born between 1970-2017 with a combination of both EA and IHPS in history. Patients were included and analyzed after parental informed consent

451

452 SNP-array analysis

453 Micro-array analysis was performed using the single-nucleotide polymorphism (SNP) CytoSNP-454 850Kv0 BeadChip (Illumina Inc., San Diego) using standard protocols and the GenomeStudio 455 genotyping module (v1.9.4, <u>www.illumnia.com</u>). Visualization of Copy Number Variations (CNVs), 456 Runs of Homozygosity (ROH) and comparisons to in-house control cohorts as well as published 457 cohorts of affected and control individuals was done using Biodiscovery Nexus CN7.5. (Biodiscovery 458 Inc., Hawthorne, CA, USA) and described previously (Brosens et al., 2016b).

459

460 Variant pre-filtering and prioritization

The initial variant filtering method has been described previously (Halim et al., 2017). In brief, we included all variants with an allele frequency below 1% in 1000 Genomes phase 3 version 5, Exome Variant Server 6500 v0.0.30, Genome of the Netherlands (Genome of the Netherlands, 2014), ExAC 0.3 and our in-house cohort (n=906), consisting of individuals captured with the SureSelect Human All Exon 50 Mb Targeted exome enrichment kit v4 (n=279), SureSelect Clinical Research Exome v1 466 (n=387) and Haloplex Exome target enrichment system (n=240), Agilent Technologies, Inc., Santa467 Clara, California).

All nonsense variants, variants predicted to affect splicing and all variants with a Combined Annotation-Dependent Depletion (CADD) score (Kircher et al., 2014) above 20 were selected for individual patient analysis in downstream tools. Different downstream tools were used to prioritize the variants. Prioritized variants were further classified according to the criteria in Table S8. Determination of variant segregation and confirmation of *de novo* of inherited status of variants was done with Sanger sequencing unless otherwise indicated.

474

475 Variant burden test and prioritization using Opal

476 We used the Variant Annotation, Analysis & Search Tool (VAAST) (Yandell et al., 2011, Hu et al., 2013, 477 Kennedy et al., 2014) cohort analysis embedded in Opal 4.29.5 (Fabric Genomics, Oakland, CA, USA) 478 to rank the variants in the individual patients. Secondly, we performed a burden test on the full 479 exomes using Exome Variant Server 6500 v0.0.30 and 1000 Genomes phase 3 version 5 as a control 480 cohort. We used a 1% allele frequency cut-off for recessive (hemizygous and homozygous) variants 481 and 0.1% cut-off for heterozygous variants. Compound heterozygosity was not considered in this 482 analysis as we did not know the phase of the haplotypes. Only putative protein changing (nonsense, 483 missense, initiator codon variants, in-frame indels, splice sites and splice regions) variants were taken 484 into account. Since we were only interested in putative deleterious variants we used an Omicia score 485 of 0.79 as a threshold as this cut-off has a false positive rate of 5%. Omicia is an algorithm included in 486 the Opal software that combines SIFT (Ng and Henikoff, 2001), PolyPhen (Adzhubei et al., 2010), 487 MutationTaster (Schwarz et al., 2010) and PhyloP (Siepel et al., 2005) to predict deleteriousness of 488 variants.

489 For the VAAST burden test we used a minimum significance of 0.05 and a gene had to have at 490 least two distinct variants in the case set. These genes were used as a gene panel in the individual 491 patient analysis. Individual variants were prioritized before individual inspection as follows. First, all 492 recessive (X-linked and putative homozygous and compound heterozygous), putative rare (MAF 493 $\leq 0.001\%$) and damaging *de novo* variants were selected. Secondly, the top 10 of variants ranked by 494 the VAAST 1.1 prioritization algorithm and subsequently the top 10 variants re-ranked by the Phevor 495 algorithm (Singleton et al., 2014) were included. We used the Human Phenotype Ontology (HPO) 496 (Singleton et al., 2014) terms esophageal atresia and pyloric stenosis as phenotype terms in the 497 algorithm. Finally, variants passing the pre-filtering criteria in genes from the burden test were 498 included.

- 499
- 500

501 Variant prioritization using bioinformatic genotype-phenotype correlation tools

502Three modules were used: PhenIX (Zemojtel et al., 2014) (http://compbio.charite.de/PhenIX/), the503Exomiser (Robinson et al., 2014) (http://www.sanger.ac.uk/resources/software/exomiser/submit/)

and the HPO prioritization incorporated within the Cartagenia software. Settings were as followed.

505 Using PhenIX the full patient phenotype in HPO terms was used, the exome target region filter is 506 on and allele frequency filter of 0.1%, pathogenicity filter was on and mode of inheritance unknown. 507 Genes were prioritized using PhenIX which compares patient phenotypes against human phenotypes 508 only. As a cut-of we used a gene relevance score of 0.8 in combination with a variant score of 0.8, or 509 a total score of 0.9.

510 When using the Exomiser tool we used similar settings: full patient phenotype in HPO terms, 511 exome target region filter is off, allele frequency filter 0.1%, pathogenicity filter on. We did not 512 remove dbSNP variants nor used an inheritance model. Genes are now prioritized using hiPhive, 513 which compares phenotypes against all species. As a cut-of we used a phenotype score of 0.8 in 514 combination with a variant score of 0.8, or an Exomiser score of 0.9.

515

516 Pathway enrichment analysis of genes affected by rare variants

To investigate if specific pathways are enriched with ultra-rare variants, Gene IDs with variants in canonical splice sites (n=16), nonsense variants (n=21), protein altering inframe InDels (n=28) and missense variants (n=557) were uploaded to Ingenuity pathway Analysis (Qiagen, Venlo, The Netherlands). Additionally, a more stringent set was uploaded with loss of function variants, predicted to be loss of function intolerant (PLI \ge 0.9, n=4) and protein altering variants with a Z-score \ge 3 (n=44).

523

524 Expression of candidate genes

525 Candidate gene expression was determined at relevant developmental time points in human and 526 mouse. Gene expression of top-ranking genes derived from the burden analysis and individual 527 patient sample prioritizations were determined using datasets (GSE13040, GSE19873, GSE34278, 528 GSE15872, GSE43381) downloaded from the Gene Expression Omnibus (GEO) (Edgar et al., 2002). 529 We used public data on mice on the endoderm, mesoderm and ectoderm at E8.25, foregut at E8.5 530 and esophagus, stomach, pyloric sphincter and intestine at E11.5-E18.5 531 (https://www.ncbi.nlm.nih.gov/geo/) (Stephens et al., 2013, Li et al., 2009, Sherwood et al., 2009, 532 Millien et al., 2008, Chen et al., 2012). These datasets were imported into BRB-ArrayTools Version: 533 4.5.0 - Beta 2. (http://linus.nci.nih.gov/BRB-ArrayTools.html), annotated by Bioconductor 534 (www.bioconductor.org), R version 3.2.2 Patched (2015-09-12 r69372) and normalized. We

determined differential expression between tissue types and classified upregulated genes beingexpressed in the tissue under investigation.

537

538 Detection of common SNP associated with IHPS

539 Genome-wide association studies (GWAS) revealed five loci highly associated with IHPS (rs11712066,

540 rs573872, rs29784, rs1933683 and rs6736913), pointing towards *MBNL1*, *NKX2-5*, *BARX1* and *EML4*

541 as candidate genes (Feenstra et al., 2012, Everett and Chung, 2013, Fadista et al., 2019). Since

- 542 rs6736913 is a low frequency missense variant, we did not further analyze this SNP in our patients.
- 543 We used Sanger sequencing to determine the risk allele frequency of the other four SNPs. With a chi
- 544 square test we compared the allele frequency in our patients with the gnomAD dataset
- 545 (<u>http://gnomad.broadinstitute.org/</u>).
- 546

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552 COMPETING INTERESTS

553 Authors do not have any potential conflicts (financial, professional, or personal) relevant to the 554 manuscript to disclose.

555

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559

560 DATA AVAILABILITY

561 Variants included in Table S4 are submitted to the ClinVar database. 562 (<u>http://www.ncbi.nlm.nih.gov/clinvar/</u>)

563

564 AUTHOR CONTRIBUTIONS STATEMENT

Conceptualization: D.T., R.W., R.H., A.K., E.B.; Methodology: R.B., H.E., W.IJ., A.K., E.B.; Software:
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A.B., H.E., H.IJ.; Data curation: E.B.; Writing – original draft: C.K., R.H., E.B.; Writing – review &
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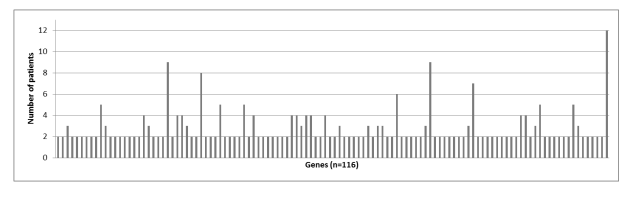
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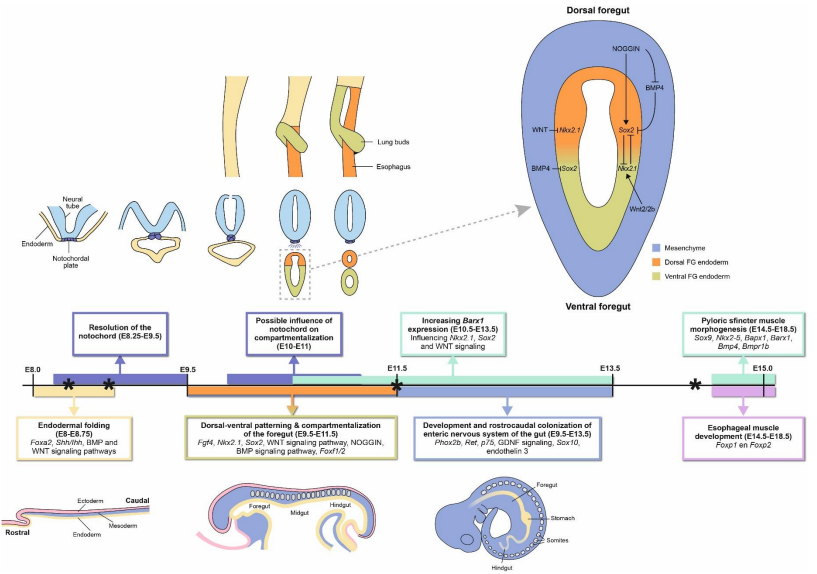


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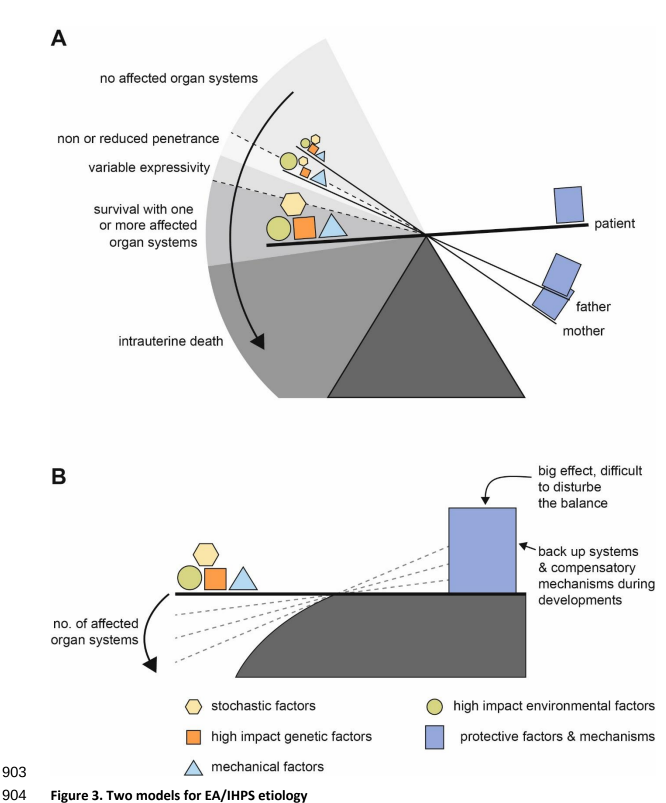
897 Figure 1. Number of patients with variants per gene

898 36 genes were found in ≥3 patients of which six genes were present in more than five patients (*CNTN2*, *DSPP*, 899 *NOTCH4*, *PRRC2A*, *SEC16B*, *ZNF717*). See also Table S6.

29



- 900 901
- Figure 2. Timeline of models and genes known to be important for foregut development in mice
- 902 (Heath, 2010, Fausett and Klingensmith, 2012, Perin et al., 2017, Anderson et al., 2006), * = time points used in expression analysis (see Table S5)



A = burden model, B = slippery slope model. The combination of multiple high impact factors (genetic, environmental, mechanical and/or stochastic) together can modulate the phenotypical spectrum. These risk factors are in balance with protective factors like backup systems and compensatory mechanisms.

Table 1. Genetic syndromes and mutated genes with tracheoesophageal and pyloric anomalies as variable features

Syndrome	Esophageal or pyloric anomaly	Inheritance	Loci	Gene(s)	OMIM	Ref
Esophageal atresia or stenosis						
Epidermolysis bullosa, junctional,	Esophageal and pyloric atresia or	AR	2q31.1	ITGA4	226730	(Varki et al., 2006, Vivona et al., 1987)
with pyloric stenosis or atresia ^c	stenosis		17q25.1	ITGA6	226730	(Ruzzi et al., 1997)
Ehlers-Danlos syndrome ^c	EA and IHPS	AD	2q32.2	COL3A1	130050	(Kroes et al., 2003, Kuivaniemi et al., 1990)
Trisomy 13	EA/TEF and IHPS	AD	13	multiple	NA	(Brosens et al., 2014, Taylor, 1968)
Trisomy 18	EA/TEF and IHPS	AD	18	multiple	NA	(Brosens et al., 2014, Taylor, 1968)
Trisomy 21	EA/TEF and IHPS	AD	21	multiple	190685	(Brosens et al., 2014, Freeman et al., 2009)
Fryns syndrome	EA/TEF and IHPS	U	unknown	unknown	229850	(Ayme et al., 1989)
Fetal alcohol syndrome	EA/TEF and IHPS	NA	NA	NA	NA	(Brosens et al., 2014, Lodha et al., 2005, Mangyanda et al., 1998)
Motility anomalies of the esophagus						
Epidermolysis bullosa dystrophia ^c	Esophageal strictures and	AR, AD	3p21.31	COL7A1	131750	(Christiano et al., 1995, Hovnanian et al., 1994,
	stenosis	AR	11q22.2	MMP1	226600	Christiano et al., 1996)
Cornelia de Lange syndrome ^{B,C}	Esophageal stenosis and	AD	5p13.2	NIPBL	122470	(Cates et al., 1989, Gillis et al., 2004)
_	dysmotility and IHPS					
Apert syndrome	Esophageal stenosis and IHPS	AD	10q26.13	FGFR2	101200	(Pelz et al., 1994, Blank, 1960)
Congenital generalized lipodystrophy	Esophageal dysmotility and IHPS	AR	17q21.2	PTRF	613327	(Rajab et al., 2010, Rajab et al., 2002)
Opitz-Kaveggia syndrome	Nutcracker esophagus and IHPS	XL	Xq13	MED12	305450	(Smith et al., 2000b, Battaglia et al., 2006)
Noonan syndrome ^c	Esophageal dysmotility and IHPS	AD	12q24.13	PTPN11	163950	(Shah et al., 1999, Barberia Leache et al., 2003)
Visceral neuropathy	Dilated non-peristaltic esophagus and IHPS	U	unknown	unknown	243180	(Schuffler et al., 1978, Tanner et al., 1976)
Costello syndrome	Loss of elastic fibers in esophagus, IHPS	AD	11p15.5	HRAS	218040	(Mori et al., 1996, Gripp and Lin, 1993)
Other associations						
Chronic idiopathic intestinal pseudo obstruction ^{B, C}	Gastro-intestinal dysmotility and IHPS	XL	Xq28	FLNA	300048	(Tanner et al., 1976, Gargiulo et al., 2007)
Fronto-metaphyseal dysplasia ^B	EA/TEF	XL	Xq28	FLNA	305620	(Franceschini et al., 1997)
X-linked periventricular heterotopia ^B	IHPS	XL	Xq28	FLNA	300049	(Nezelof et al., 1976)
FG syndrome ^{B, C}	Esophageal dysmotility and IHPS	XL	Xq28	FLNA	300321	(Unger et al., 2007, Peeters et al., 2012)
CHARGE syndrome ^{B, C}	EA/TEF	AD	8q12.1-	CHD7	214800	(Brosens et al., 2014)
	,	-	q12.2			· ···· · · · · · · · · · · · · · · · ·
Hypogonadotropic hypogonadism with or without anosmia ^{B, C}	IHPS ^A	AD	8q12.1- q12.2	CHD7	612370	(Jongmans et al., 2009, Kim et al., 2008)

- 910 This table is modified from two reviews on esophageal atresia (Brosens et al., 2014) and infantile hypertrophic pyloric stenosis (Peeters et al., 2012). AD; autosomal dominant, AR;
- 911 autosomal recessive, U; unknown, NA; not applicable, XL; X-linked; EA; esophageal atresia, TEF; trachea-esophageal fistula, IHPS; infantile hypertrophic pyloric stenosis. ^A In literature
- 912 IHPS is associated with other genes responsible for this syndrome. ^B No overlap in EA and IHPS phenotype for this syndrome, the gene mutated in this syndrome can be responsible
- 913 for different syndromes in which either EA or IHPS are variable features. ^c More genes associated to possible several subtypes of this syndrome.

914 Table 2. Phenotype description

915

Individual	Gender	EA-	Phenotype	Remarks
		type		
SKZ_0027	female	С	EA/TEF, IHPS, thin ear helix, seizures	-
SKZ_0096	male	С	EA/TEF, IHPS, syndactyly second-third finger, radial dysplasia, abnormal fibula	VACTERL association
SKZ_0244	male	С	EA/TEF, IHPS, anal atresia, intestinal malrotation, sacral dimple, abnormal os coccygis, abnormal vertebrae L1, thenar hypoplasia, both sides hypoplastic "floating" thumbs, both sides dysplastic radii	VACTERL association, mother is a DES daughter
SKZ_0321	male	С	EA/TEF, IHPS, mild left sided expansion of the pyelocaliceal system , breath holding spells	-
SKZ_0353	female	С	EA/TEF, IHPS, sacral dimple, thin/slender build, diminished hearing, palpebral fissures slant up, hemolytic anemia, short phalanges	Glucose-6-phosphate dehydrogenase deficiency
SKZ_0399	male	С	EA/TEF, IHPS, anal atresia, sacral dimple, 2 umbilical vessels, posteriorly rotated ears, small ears/microtia, flat face, bifid scrotum, small penis/micropenis, small palmar crease, thick fingers, broad thumbs, proximal placement of thumbs, microstomia, thick broad neck, wide nasal bridge, patent ductus arteriosis, 4 th toe abnormally placed	VACTERL association
SKZ_0400	male	С	EA/TEF, IHPS, extra ribs, fusion of vertebrae, macrocephaly, bulbar dermoid cyst, auricular tags, short thick/broad neck	Klippel-Feil syndrome
SKZ_0683	male	С	EA/TEF, IHPS, sacral dimple	-
SKZ_0760	male	С	EA/TEF, IHPS, hemivertebrae, bitemporal narrowing of the head, prominent forehead, hyper mobile/ extensible fingers, narrow thorax/funnel chest, thin lower and upper lip, spasticity, cerebral palsy	-
SKZ_0788	male	С	EA/TEF, IHPS, inguinal hernia, jaundice, deafness	-
SKZ_0790	female	С	EA/TEF, IHPS	-
SKZ_0796	male	С	EA/TEF, IHPS	Vanishing twin
SKZ_0848	male	С	EA/TEF, IHPS, sacral dimple, hypospadias, patent ductus arteriosus	-
SKZ_0887	male	С	EA/TEF, IHPS, abnormal sacrum, fusion of vertebrae, posteriorly rotated ears, small mandible/micrognathia, rocker- bottom feet, sandal gap of toes, open mouth appearance, short neck, jaundice	-
SKZ_1003	male	С	EA/TEF, IHPS, abnormal sacrum, cleft jaw, cleft palate, cleft upper lip, depressed/flat nasal bridge, fused ribs	Methyldopa (aldomet) for hypertension during pregnancy
SKZ_1248	female	С	EA/TEF, IHPS, small large fontanel, deafness, small ears, auricular tags, single palmar crease, small/hypoplastic deep set ears	
SKZ_1260	male	С	EA/TEF, IHPS, syndactyly of second-third toe, bifid/fused ribs	-
SKZ_1353	male	С	EA/TEF, IHPS, cleft uvula, epicanthic folds, abnormal dermatoglypic patterns, hyperconvex/clubbed nails, hypoplastic scrotum, hypospadias, bifid scrotum, hydrocele of testis	-
SKZ_1407	female	А	EA, IHPS	-
SKZ_1472	male	С	EA/TEF, IHPS, eczema of hands with hyperhidrosis, blisters and erythema, Xerosis Cutis	Antibiotics for respiratory infection during pregnance
SKZ_1961	male	С	EA/TEF, IHPS, sacral dimple, mild dysmorphic features, small mouth, pointy ears, long fingers	Maternal hypertension
SKZ_2013	male	А	EA, IHPS, persistent superior vena cava, scoliosis, Horner's syndrome	-
SKZ_2023	male	С	EA/TEF, IHPS, small chin, sacral dimple	-
SKZ_2050	male	С	EA/TEF, IHPS, atrial septum defect	
SKZ_2082	male	С	EA/TEF, IHPS, persistent tracheolaryngeal cleft, anal atresia, atrial septum defect, tracheal-laryngeal anomaly, prostate fistula	VACTERL association
SKZ_2149	male	С	EA/TEF, IHPS	-
SKZ_2171	female	С	EA/TEF, IHPS, spina bifida Th10/11, synostoses vertebrae, hydronephrosis, kyphoscoliosis	Unknown medication for headaches and nerves during pregnancy

EA; esophageal atresia, TEF; trachea-esophageal fistula, IHPS; infantile pyloric stenosis, DES; di-ethylstilbestrol. EA-type classification according to Gross classification (Gross, 1947)

919

Gene	Chr.	Start position	rsID	HGVS c	VAAST p-value
ALMS1	chr2	73828337	rs201777220	c.11885T>C	1,00E-06
ALMS1	chr2	73746907		c.9543delA	1,00E-06
SLC28A3	chr9	86895771	rs762396296	c.1673dupA	0,002853
SLC28A3	chr9	86912927	rs775359011	c.677G>A	0,002853
SP2	chr17	46002296	rs778698435	c.1384C>A	0,002853
SP2	chr17	45993628	rs754352270	c.191C>T	0,002853
EPB41	chr1	29435944		c.2410A>G	0,019802
EPB41	chr1	29391549	rs768609152	c.2063T>G	0,019802
AMBRA1	chr11	46563580	rs755884183	c.1717C>G	0,019802
AMBRA1	chr11	46567274		c.431C>T	0,019802
VWA8	chr13	42293773	rs371770462	c.3070G>A	0,019934
VWA8	chr13	42439870	rs201163045	c.1425+2T>C	0,019934
CLGN	chr4	141320158	rs201306926	c.731A>G	0,039604
CLGN	chr4	141315036	rs200652126	c.1309G>T	0,039604
SDK2	chr17	71357964	rs147983543	c.5326G>A	0,049505
SDK2	chr17	71415318		c.2173G>A	0,049505
PDLIM7	chr5	176911101	rs200609502	c.1141A>G	0,049505
PDLIM7	chr5	176916502	rs764108486	c.761C>T	0,049505
GUCY2F	chrX	108652306	rs7883913	c.1883G>A	0,019802
GUCY2F	chrX	108638614	rs35726803	c.2380G>A	0,019802

N/A = not applicable

922 Table 4. Summary of overlapping top candidate genes

		EA/IHPS pa	tients	(n=15)		Healthy controls (n=44)			
		are (MAF ≦0.001%)	Ultra-rare (MAF 0%)		Rare (MAF ≤0.001%)		Ultra-rare (MAF 0%)		
	LB	VUS/LD	LB	VUS/LD	LB	VUS/LD	LB	VUS/LD	
Important in normal foregut development (Fig. 2)	1	1	0	0	3	5	1	2	
Genes associated with genetic syndromes involving both EA and IHPS as variable features (Table 1)	1	0	0	0	2	4	1	1	
Genes associated with IHPS (Peeters et al., 2012)	0	0	0	0	0	2	0	0	
Genes involved in neuromuscular and connective tissue syndromes associated with IHPS (Peeters et al., 2012)	1	1	1	0	1	2	1	0	
Genes involved in syndromes and signaling disturbances associated with IHPS (Peeters et al., 2012)	0	3	0	0	1	6	1	1	
Genes involved in ciliopathies and disturbances of gene regulation associated with IHPS (Peeters et al., 2012)	0	1	0	0	4	2	1	0	
Genes involved in lymphatic abnormalities and syndromes of environmental and unknown origin associated with IHPS (Peeters et al., 2012)	0	0	0	0	1	0	0	0	
Genes involved in genetic syndromes and abnormalities	1	1	1	1	6	6	0	1	

923 Variant allele count per gene. LB = likely benign, VUS = variant of unknown significance, LD = likely deleterious.

924 See Table S6 for the complete results, adapted from Peeters et al. en Brosens et al. (Peeters et al., 2012, Brosens et 925 al., 2014).

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928 Table 5. Comparison with control cohort: number of variants

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		Ultra-rare vai	riants (MAF 0	%)	Rare variants (MAF ≤0.001%)				
	EA/IHPS patients (n=15)		Healthy controls (n=44)		EA/IHPS patients (n=15)		Healthy control (n=44)		
		#		#		#		#	
Putative deleterious	296	19.73	725	16.48	912	60.8	2667	60.6	
LOF intolerant	28	1.87	85	1.93	81	5.4	272	6.2	
De novo variants	0	-	Unknown	-	0	-	Unknown	-	
Recessive	291	19.4	715	16.25	898	59.9	2631	59.80	
Compound heterozygous	3	0.2	3	0.06	8	0.53	6	0.14	
X-linked	11	0.73	23	0.52	28	1.87	95	2.16	

930 # = number of variants divided by the number of patients. A Chi² test showed no significant

931 differences between the patients and the healthy controls (all p>0.05)

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