Vistara non-invasive prenatal screen



Vistara identifies probability for conditions that may have otherwise gone undetected until after birth or into childhood. All conditions are inherited in an autosomal or X-linked dominant fashion, which means that if the mutation is present, the child will be affected by the condition and experience related symptoms.

Condition ¹ <i>Gene(s)</i>	Clinical synopsis ^{2,3}	Cases caused by de novo mutations ^{2,3}	Ultrasound findings ^{2,3}			Clinical	Detection
			None	Third trimester	Non- specific	actionability	rate for gene ¹
Achondroplasia FGFR3	The most common form of skeletal dysplasia; may cause hydrocephalus, delayed motor milestones, and spinal stenosis	80%		•	•	Labor and delivery management, monitor for spinal stenosis, early sleep studies to reduce risk of SIDS	>96%
Alagille syndrome JAG1	Affects multiple organ systems and may cause growth problems, congenital heart defects, and vertebral differences	50% to 70%	•			Symptom-based treatment	>86%
Antley Bixler syndrome FGFR2	A type of craniosynostosis; also causes premature fusion of the arm bones, blockage of the nasal passage, and permanently flexed or extended joints	more severe forms		•		Fetal MRI, avoid instrumented delivery, corrective surgery, monitor for hydrocephalus	>96%
Apert syndrome FGFR2	A type of craniosynostosis; also causes abnormal formation of the fingers, toes, and vertebrae, and other organ anomalies	more severe forms		•		Fetal MRI, avoid instrumented delivery, corrective surgery, monitor for hydrocephalus	>96%
Cardiofaciocu- taneous syndrome 1,3,4 BRAF, MAP2K1, MAP2K2	Causes abnormalities of the heart, face, skin, and hair; may cause developmental delays and intellectual disability	majority		•	•	Fetal echocardiogram	>96%
CATSHL syndrome FGFR3	Acronym stands for camptodactyly, tall stature, scoliosis, and hearing loss; may increase risk for intellectual disability	unknown	•			Early adoptionof sign language and behavioral intervention	>96%
CHARGE syndrome CHD7	Acronym stands for coloboma, heart defects, atresia of the choanae, retardation of growth and development, genital abnormality, ear abnormalities; may cause hearing loss, developmental delays, and cleft lip and/or palate	majority		•	•	Early referral to endocrinology, adoption of sign language, and behavioral intervention	>91%
Cornelia de Lange syndrome 1,2,3,4,5 NIPBL, SMC1A, SMC3, RAD21, HDAC8	Causes a range of physical, cognitive, and medical challenges	99%	•		•	Monitor for cardiac, GI, and limb comorbidities	53% to >96%
Costello syndrome HRAS	Causes heart defects, intellectual disability, developmental delays, growth delays, and increased risk of malignant tumors	majority	•		•	Nasogastric or gastronomy feeding, behavioral and medical intervention	>92%
Crouzon syndrome FGFR2, FGFR3	A type of craniosynostosis; also causes hearing loss and dental problems in some cases	more severe forms		•		Fetal MRI, avoid instru- mented delivery, corrective surgery, monitor for hydrocephalus, early adoption of sign language	>96%
Ehlers-Danlos syndrome, classic, type VIIA, cardiac valvular form, type VIIB COL1A1, COL1A2	Causes defects in connective tissue that can vary from mildly loose joints to life-threatening complications, such as aortic dissection	50%	•			Orthotic treatment, monitoring for vascular complications	>92%
Epileptic encepha- lopathy, early infantile, 2	Causes seizures with secondary developmental delay	majority	•			Monitor and treat seizures	>84%

Condition ¹ Gene(s)	Clinical synopsis ^{2,3}	Cases caused by de novo mutations ^{2,3}	Ultra	sound findi	ngs ^{2,3}	Clinical actionability	Detection rate for gene ¹
			None	Third trimester	Non- specific		
Hypochondro- blasia FGFR3	Causes a mild form of dwarfism; may cause seizures with secondary developmental delay	up to 80%	•			Monitor and treat seizures	>96%
ntellectual disability SYNGAP1	Causes intellectual disability and developmental delays	~100%	•			Early behavioral interventions	>86%
Jackson Weiss syndrome FGFR2	A type of craniosynostosis; also causes foot abnormalities	more severe forms		•		Fetal MRI, avoid instrumented delivery, corrective surgery, monitor for hydrocephalus	>96%
Juvenile myelomonocytic eukemia (JMML) PTPN11	A rare pediatric blood cancer; five- year survival is approximately 50%	unknown	•			Monitor bloodwork and medical intervention	>96%
ECPARD syndrome 1,2 Noonan syndrome with multiple entigines) PTPN11, RAF1	Similar to Noonan syndrome, with notable brown skin spots (lentigines); causes short stature, heart defects, bleeding problems, and, in some cases, mild intellectual disabilities	unknown	•		•	Fetal echocardiogram	>96%
Muenke syndrome FGFR3	A type of craniosynostosis; may cause hearing loss, developmental delays, and cleft lip and/or palate	unknown		•		Fetal MRI, corrective surgery, early adoption of sign language, and behavioral intervention	>96%
Noonan syndrome 1,3,4,5,6,8,9 PTPN11, SOS1, RAF1, RIT1, KRAS, NRAS, SOS2, BHOC2, BRAF, MAP2K1, HRAS, CBL	Causes short stature, heart defects, bleeding problems, and, in some cases, mild intellectual disabilities	25% to 70%	•	•	•	Fetal echocardiogram, labor and delivery management, early assessment for learning differences	>92% to >96%
Osteogenesis mperfecta, cype I,II,III,IV COL1A1, COL1A2	Causes fragile bones that break easily, often without an identifiable cause	more severe forms	•	•		Labor and delivery management, neonatal care, early recognition and treatment of fractures	>92%
Pfeiffer syndrome type 1,2,3 FGFR2	A type of craniosynostosis; also causes hearing loss, intellectual disability, hand abnormalities, and may result in early death	more severe forms		•		Fetal MRI, avoid instrumented delivery, corrective surgery, monitor for hydrocephalus, early adoption of sign language, and behavioral intervention	>96%
Rett syndrome MECP2	Causes a rapid regression in language and motor skills at 6 to 18 months of age; autism, seizures, and long QT syndrome are often present	>99%	•			Evaluate for cardiac risk, monitor and treat seizures, early medical and behavioral interventions	>78%
Sotos syndrome 1 NSD1	Overgrowth syndrome; also causes developmental delays, intellectual disability, and behavioral problems	>95%		•		Fetal echocardiogram, fetal renal ultrasound, and early behavioral intervention	>47%
Thanatophoric dysplasia, dypes I,II -GFR3	A severe skeletal disorder that typically results in stillbirth or neonatal death due to respiratory failure	majority			•	Labor and delivery management	>96%
Fuberous sclerosis 1,2 ISC1, TSC2	Causes benign tumor growth in many organ systems in the body that can be life-threatening; may also cause seizures and secondary developmental delays	66%				Fetal echocardiogram, postnatal MRI, medical and behavioral interventions	>91% to >96%

= some types or cases

- References
 1. Validation data, Baylor. 2020.
 2. GeneReviews. https://www.ncbi.nlm.nih.gov/books/NBK1116/
 3. Genetics Home Reference. https://ghr.nlm.nih.gov/

Vistara has been developed and its performance characteristics determined by the CLIA-certified laboratory performing the test. The test has not been cleared or approved by the US Food and Drug Administration (FDA). CAP accredited, ISO 13485 certified, and CLIA certified. © 2021 Natera, Inc. All Rights Reserved.