

## THIS IS NOT A TEST REQUEST FORM. Please complete and submit with the test request form or electronic packing list.

## **GENOMIC MICROARRAY PATIENT HISTORY FORM**

Sex Assigned at Birth: □Female □Male □Intersex Ordering Provider: Practice Specialty:		Gender Identity (optional):					
				Clinical Information (check all t	hat apply)		
				Perinatal History	Behavioral	Craniofacial	Gastrointestinal
				Not evaluated	Not evaluated	Not evaluated	Not evaluated
				Within normal limits	🗆 Within normal limits	Within normal limits	Within normal limits
Prematurity	Autism spectrum disorder	🗆 Cleft lip +/- cleft palate	🗆 Gastroschisis				
	Oppositional defiant disorder	🗆 Cleft palate alone	Hirschsprung disease				
Oligohydramnios	□ Obsessive compulsive disorder	🗆 Coloboma	Omphalocele				
Polyhydramnios	Pervasive developmental delay	🗆 Craniosynostosis	Pyloric stenosis				
Other:	□ Other	Dysmorphic facial features	🗆 Tracheoesophageal fistula				
Growth	Neurological	Ear malformation	Other:				
□ Not evaluated	□ Not evaluated		Genitourinary				
Within normal limits	Within normal limits	□ Microcephaly	□ Not evaluated				
Failure to thrive	Ataxia/dystonia/chorea	List HC, if known:	Within normal limits				
Overgrowth	□ Hypotonia	Other:	🗆 Ambiguous genitalia				
□ Short stature	□ Neural tube defect	Cutaneous	□ Hydronephrosis				
Other.	□ Seizures	□ Not evaluated	□ Hypospadias				
	Spasticity	🗆 Within normal limits	□ Kidney malformation				
Development	Structural brain anomaly	Hyperpigmentation	Undescended testis				
□ Within normal limits	Other:	Hypopigmentation	Urethra/ureter obstruction				
□ Fine motor delay	Cardiac	Musculoskeletal	Other:				
□ Gross motor delay	□ Not evaluated	□ Not evaluated	Family History				
□ Speech delay	Within normal limits	Within normal limits	□ Not evaluated				
Other.	□ ASD	Contractures	No relevant family history				
	□ AV canal defect	□ Club foot	Parents with				
Cognitive	Coarctation of aorta	🗆 Diaphragmatic hernia	≥ two miscarriages				
<ul> <li>Not evaluated</li> <li>Within normal limits</li> </ul>	Hypoplastic left heart	□ Limb anomaly	□ Other relatives with				
	Tetralogy of Fallot	□ Polydactyly	similar clinical history				
Learning disability     Intellectual disability		□ Scoliosis	(explain below)				
Intellectual disability I List IQ/DQ, if known: I Other:	Other.	Syndactyly	(p				
	-	Vertebral anomaly					
	_	Other:					

Clinical Descriptions-Include any additional relevant clinical information not provided above.

In cooperation with the National Institutes of Health's effort to improve understanding of specific genetic variants, ARUP submits HIPAA-compliant, deidentified (cannot be traced back to the patient) genetic test results and health information to public databases. The confidentiality of each sample is maintained. If you prefer that your test result not be shared, call ARUP at 800-242-2787 ext. 3301. Your deidentified information will not be disclosed to public databases after your request is received, but a separate request is required for each genetic test. Additionally, patients have the opportunity to participate in patient registries and research. To learn more, visit aruplab.com/genetics.

For questions, contact an ARUP genetic counselor at 800-242-2787 ext. 2141

Master Label