



Chromosome	Start (bp)	Stop (bp)	Gene	Band	Syndrome	OMIM	ISCA 4x44k	ISCA & ISCA UPD 4x180k	ISCA 8x60k	Syndrome Plus 2x105k
chr10	87349292	88116230	GRID1	10q22	10q22-23 Deletion		•	•	•	#
chr10	83625077	84735341	NRG3	10q22	10q22-23 Deletion		•	•	•	#
chr11	8634369	86344081	FZD4	11q14	11q14 Microdeletion		•	•	•	#
chr11	114500000	130800000		11q23.3-q24.3	11q23.3 Jacobsen deletion	147791				#
chr12	65029066	65359020	GRIP1	12q14	12q14.3 Deletion		•	•	•	#
chr13	47300000	59600000		13q14.2-q21.1	13q14 Deletion		•	•	•	#
chr14	60182506	60185933	SIX1	14q22	14q22 Microdeletion		•	•	•	#
chr14	60246009	60260792	SIX4	14q22	14q22 Microdeletion		•	•	•	#
chr14	60045150	60048904	SIX6	14q22	14q22 Microdeletion		•	•	•	#
chr14	56337178	56346937	OTX2	14q22	14q22 Microdeletion, microphthalmia, syndromic 5		•	•	•	#
chr14	53486207	53493362	BMP4	14q22	14q22 Microdeletion, Orofacial Cleft 11		•	•	•	#
chr15	30110018	30248527	CHRNA7	15q13	15q13.3 Microdeletion	612001	•	•	•	#
chr15	41612952	41664382	KIAA0377 (HISPPD2A)	15q15.3	15q15.3 Infertility and deafness		•	•	•	#
chr15	72700000	75700000		15q24.1	15q24.1 Microdeletion		•	•	•	#
chr16	28100000	34600000		16p11.2	16p11.2 Deletion	611913	•	•	•	#
chr16	21200000	34600000		16p11.2-p12.2	16p11.2-p12.2 Microdeletion		•	•	•	#
chr16	14800000	16800000		16p13.1	16p13.1 Microdeletion predisposition to autism		•	•	•	#
chr17	41217449	41289373	CRHR1	17q21.31	17q21.31 Microdeletion	610443	•	•	•	#
chr17	41327544	41461546	MAPT	17q21.31	17q21.31 Microduplication	610443	•	•	•	#
chr18	70416108	70906616	ZNF407	18q22	18q Deletion	301808	•	•	•	#
chr1	2149994	2231416	SKI	1p36	1p36 Microdeletion	607872	•	•	•	#
chr1	3548988	3646715	TP73	1p36	1p36 Microdeletion	607872	•	•	•	#
chr22	21852552	21990224	BCR	22q11	22q11.2 Distal deletion	611867	•	•	•	#
chr22	17900000	25900000		22q11.2	22q11.2 Microduplication	608363	•	•	•	#
chr22	49400000	51300000	SHANK3	22q13.33	22q13.33 Microdeletion (Phelan McDermid)	606232	•	•	•	#
chr2	61136239	61673319	MicroDeletionRegion	2p15	2p15-p16.1 Microdeletion		•	•	•	#
chr2	239634801	239987580	HDAC4	2q37	2q37.3 Monosomy		•	•	•	#
chr3	197951312	198043751	PAK2	3q29	3q29 Microdeletion	609425	•	•	•	#
chr6	1555680	1559128	FKHL7(FOXC1)	6p24	6p24 Deletion	612852	•	•	•	#
chr6	155500001	161000000		6p25.3	6p25.3 Microdeletion	220210	•	•	•	#
chr8	11571877	11654918		8p23.1	8p23.1 Microduplication		•	•	•	#
chr9	831690	959090	Dmental RetardationT1	9p24	9p Deletion	158170	•	•	•	#
chr9	1039858	1047552	Dmental RetardationT2	9p24	9p Deletion	158170	•	•	•	#
chr9	140000000	140000000	EHMT1	9q34.3	9q34.3 Microdeletion	610253	•	•	•	#
chrX	54488612	54539324	FGD1	Xp11	Aarskog-Scott	305400	•	•	•	#
chr15	98500000	102531392		15q26.3	Abnormal growth		•	•	•	#
chrX	30232507	30237413	NR0B1(DAX1)	Xp21	Adrenal hypoplasia congenita	300200	•	•	•	#
chrX	152643530	152663374	ABCD1	Xq28	Adrenoleukodystrophy	300475	•	•	•	#
chr5	126140732	126206008	LMNB1	5q23	Adult-onset autosomal dominant leukodystrophy	169500	•	•	•	#
chrX	100491098	100532426	BTK	Xq22.1	Agammaglobulinemia, X-linked	300755	•	•	•	#
chr20	10566334	10602590	JAG1	20p12	Alagille	118450	•	•	•	#
chr20	56838189	56929641	GNAS	20q13	Albright hereditary osteodystrophy	103580	•	•	•	#
chr16	155725	171196	HBA1&HBA2	16p13	Alpha thalassemia mental retardation	141750	•	•	•	#
chrX	76647847	76928358	ATRX	Xq21	Alpha thalassemia mental retardation	301040	•	•	•	#
chrX	105769810	107827431	COL4A5	Xq22.3	Alport, X-linked	301050	•	•	•	#
chrX	66680599	66860844	AR	Xq12	Androgen insensitivity	300068	•	•	•	#
chr15	21429787	21452081	MAGEL2	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	21361547	21407800	MKRN3	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	20594720	20637877	NIPA1	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	20556790	20585849	NIPA2	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22713304	22720548	PWS_ICRegion	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22747536	22751866	PWS_ICRegion	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22768233	22793232	SNORD107/64/108	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	23064582	23084648	SNORD109B	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22835000	23010100	SnoRNA	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22819887	22774822	SNRPN	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22751228	22774822	SNRPN	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	22741227	22784821	SNURF	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr15	21471646	21493542	NDN	15q11	Angelman / Prader Willi	105830/176270	•	•	•	#
chr11	31762916	31796085	PAX6	11p13-p14	Aniridia	106210	•	•	•	#
chr1	145694956	145712108	GJA5	1q21	Atrial fibrillation		•	•	•	#
chr5	172591744	172594868	NKX2-5	5q35	Atrial septal defect with atrioventricular conduction defects	108900	•	•	•	#
chr1	229829184	230243641	DISC1	1q42	Autism	209850	•	•	•	#
chr2	50000992	51113178	NRXN1	2p16	Autism	209850	•	•	•	#
chr2	162189192	162548489	SLC4A10	2q24	Autism	209850	•	•	•	#
chr3	2115550	3074645	CNTN4	3p25	Autism	209850	•	•	•	#
chr7	121746687	122313750	CADPS2	7q31	Autism	209850	•	•	•	#
chr7	145444386	147749019	CNTNAP2	7q35-36	Autism	209850	•	•	•	#
chr7	154943160	154950579	EN2	7q36	Autism	209850	•	•	•	#
chr7	116099695	116225676	MET	7q31	Autism	209850	•	•	•	#
chr10	64241763	64246133	EGR2	10q21	Autism	209850	•	•	•	#
chr15	23463512	23669962	ATP10A	15q12	Autism	209850	•	•	•	#
chr15	24339787	24767950	GABRB3	15q12	Autism	209850	•	•	•	#
chr22	49459936	49518507	SHANK3	22q13	Autism	209850	•	•	•	#
chrX	70281436	70307776	NLGN3	Xq13	Autism	209850	•	•	•	#
chrX	5819885	61574822	NLGN4X	Xp22	Autism	209850	•	•	•	#
chrY	13052929	13603474	AZFa	Yq11.2	AZFa region	415000	•	•	•	#
chrY	23663736	25513794	AZFb	Yq11.2	AZFb region	415000	•	•	•	#
chrY	23539798	23560998	BPY2	Yq11.2	AZFb region	415000	•	•	•	#
chrY	25173539	25194740	BPY2	Yq11.2	AZFb region	415000	•	•	•	#
chrY	25586438	25607639	BPY2	Yq11.2	AZFb region	415000	•	•	•	#
chrY	24600764	24603549	CDY1	Yq11.2	AZFb region	415000	•	•	•	#
chrY	26177652	26180436	CDY1	Yq11.2	AZFb region	415000	•	•	•	#
chrY	23684890	23754627	DAZ1	Yq11.2	AZFb region	415000	•	•	•	#
chrY	24765503	24770366	GOLGA2LY(AF332229)	Yq11.2	AZFb region	415000	•	•	•	#
chrY	26010846	26015709	GOLGA2LY(AF332229)	Yq11.2	AZFb region	415000	•	•	•	#
chr15	46285791	46382417	SLC12A1	15q21	Barter 1	601678	•	•	•	#
chr11	128213125	128242478	KCNJ1	11q24	Barter 2	241200	•	•	•	#
chr1	16221073	16256063	CLCNKB	1p36	Barter 3	607364	•	•	•	#
chr1	55237205	55247053	BSND	1p32	Barter 4A	602522	•	•	•	#
chr1	16217957	16233132	CLCNKA	1p36	Barter 4B	613090	•	•	•	#
chr3	12190250	123488032	CASR	3q13.33-q21.1	Barter with autosomal dominant hypocalcaemia	601199	•	•	•	#
chr11	2851440	2873550	CDKN1C	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr11	1962981	1965640	H19	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr11	2069825	2136469	IGF2	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr11	2127584	2148999	INS	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr11	2896078	2917225	PHLDA2	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr11	2867526	2913051	SLC22A18	11p15	Beckwith-Wiedemann	130650	•	•	•	#
chr16	56211151	56256445	GPR56	16q13	Bilateral frontoparietal polymicrogyria	606854	•	•	•	#
chr3	140144730	140149880	FOXL2	3q22	Blepharophimosis	110100	•	•	•	#
chrX	133335008	133390488	PHF6	Xq26	Borjeson-forssman-lehmann	301900	•	•	•	#
chr13	90848888	92317488	GPC5	13q31	Brachydactyly		•	•	•	#
chr13	92677096	93853948	GPC6	13q31	Brachydactyly		•	•	•	#
chr20	33360416	33489441	GDF5	20q11	Brachydactyly type C	113100	•	•	•	#
chr8	72272222	72437021	EYA1	8q13	Branchio-oto-renal	113650	•	•	•	#
chr13	31787617									

chr7	91100001	92800000	PDCD10	7q21.2	Cerebral cavernous malformations (CCM)	603285					
chr17	15073822	15109369	PMP22	17p11	Charcot marie tooth disease	118220					
chrX	106758415	106780912	PRPS1	Xq22	Charcot marie tooth disease, X-linked	311070					
chr8	61753893	61942017	CHD7	8q12	CHARGE	214800					
chrX	2862852	2892311	CDPX1(ARSE)	Xp22	Chondrodysplasia punctata, X-linked recessive	302950					
chrX	85002841	85189222	CHM	Xp21	Choroideremia	303100					
chrX	37524264	37557658	CYBB	Xp11	Chronic granulomatous disease	306400					
chr5	147184339	147191453	SPINK1	5q32	Chronic pancreatitis	167800					
chr4	4911225	4917628	MSX1	4p16	Cleft lip	608874					
chr2	199842469	200038076	SATB2	2q32	Cleft pallet	119540					
chr6	45404032	45626796	RUNX2	6p21	Cleidocranial dysplasia	119600					
chrX	20077950	20194671	RPS6KA3	Xp22	Coffin-lowry	303600					
chr6	32044642	32162690	CYP21A2	6p21.32	Congenital adrenal hyperplasia (CAH)	201910					
chr15	91244423	91372241	CHD2	15q26	Congenital diaphragmatic hernia	142340					
chr15	94670161	94683047	NR2F2	15q26	Congenital diaphragmatic hernia	142340					
chr8	106400323	106885943	ZFPM2(FOG2)	8q23	Congenital heart	610187					
chr5	36912742	37101678	NIPBL	5p13	Cornelia de Lange	122470					
chrX	53467795	53486343	SMC1L1(SMC1A)	Xp11	Cornelia de Lange, X-linked	300590					
chr10	88506376	88674925	BMPRIA	10q23	Cowden	158350					
chr5	174084181	174090508	MSX2	5q35	Craniosynostosis	604757					
chr11	15949900	16454494	SOX6	11p15.1-p15.2	Craniosynostosis	128350					
chr16	34877510	35027509	Centromere	16p11	Creatine deficiency / X-linked mental retardation	300352					
chr16	33686002	33694027	SLC6A8	16p11	Creatine deficiency / X-linked mental retardation	300352					
chrX	152606586	152615234	SLC6A8	Xq28	Creatine deficiency / X-linked mental retardation	300352					
chr5	1306287	1348162	TERT	5p15	Cri-du-Chat	123450					
chr13	31211679	31275009	LGR8(RFXP2)	13q13	Cryptorchidism, unilateral or bilateral	219050					
chr7	156479506	156496108	HLXB9(MNX1)	7q36	Currarin	176450					
chr7	117120017	117308716	CFTR	7q31.2	Cystic fibrosis	219700					
chr17	3486511	3513146	CTNS	17p13	Cystinosis	219800					
chr19	33321421	33360683	SLC7A9	19q13.11	Cystinuria	220100					
chr2	44399406	44442448	PREPL	2p21	Cystinuria with mitochondrial disease	606407					
chr2	44356103	44401443	SLC3A1	2p21	Cystinuria with mitochondrial disease	606407					
chr3	148609871	148617196	ZIC1	3q24	Dandy-Walker	220200					
chr3	148596527	148607097	ZIC4	3q24	Dandy-Walker	220200					
chrX	119444035	119487189	LAMP2	Xq24	Danon disease	300257					
chrX	1.53E+08	1.53E+08	DCX	Xq22.3-q23	DCX-associated lissencephaly/subcortical band heterotopia	300067					
chr5	149717428	149760063	TCOF1	5q33	Deafness	154500					
chr13	19659605	19665114	GJB2(Connexin 26)	13q12	Deafness	220290					
chrX	100487306	100490343	TIMM8A	Xq22	Deafness-dystonia-optic neuropathy	304700					
chr16	65620551	65692459	CBFB	16q22	Delayed cranial ossification						
chrX	152821179	152825834	AVPR2	Xq28	Diabetes insipidus, nephrogenic, X-linked	304800					
chr6	144370000	144377576	HYMAI	6q24	Diabetes mellitus, transient neonatal, 1	601410					
chr6	144293129	144427428	ZAC(PLAGL1)	6q24.2	Diabetes mellitus, transient neonatal, 1	601410					
chr19	47055828	47067324	RPS19	19q13	Diamond blackfan anemia	105650					
chr8	11571877	11654918	GATA4	8p23	diGeorge 1	188400					
chr8	8679409	8788541	MFHAS1	8p23	diGeorge 1	188400					
chr22	19601714	19638035	CRKL	22q11	diGeorge 1	188400					
chr22	17698224	17815220	HIRA	22q11	diGeorge 1	188400					
chr22	18124226	18151110	TBX1	22q11	diGeorge 1	188400					
chr10	8136673	81571170	GATA3	10q14	diGeorge 2	60362					
chrX	31048667	33277779	DMD	Xp21.1-p21.2	DMD	310200					
chr14	54100000	58100000		14q22.2-q22.3	Dopa resistant dystonia / parkinsons	186600					
chr21	34810654	34909252	DSCR1(RCAN1)	21q22	Downs syndrome critical region	190685					
chr21	37517596	37561703	DSCR3	21q22	Downs syndrome critical region	190685					
chr21	37661729	37809549	DYRK1A	21q22	Downs syndrome critical region	190685					
chr21	35081968	36278917	RUNX1	21q22	Downs syndrome critical region	190685					
chr18	44824170	45250999	DYM	18q21	Dygve-delchlor-clausen disease	223800					
chr7	94113623	94146938	PEG10	7p21	Dystonia-11	159900					
chr7	94123295	94123994	PEG10_ICRRegion	7p21	Dystonia-11	159900					
chr7	94042477	94133413	SGCE	7p21	Dystonia-11	159900					
chrX	68752636	69176046	EDA	Xq13	Ectodermal dysplasia	305100					
chr7	93861809	93898480	COL1A2	7q21	Ehlers-danlos	225320					
chrX	147100000	155270560	EMD	Xq28	Emery-dreifuss muscular dystrophy	310300					
chrX	100539435	100549657	GLA	Xq22	Fabry disease	301500					
chr3	10068113	10141343	FANCD2	3p25.3	Fanconi anemia	227650					
chr16	88331480	88410566	FANCA	16q24	Fanconi anemia	227650					
chr2	15998134	16004579	MYCN	2q24	Feingold	164280					
chrX	148801201	148840303	FMR1	Xq27	Fragile X	300824					
chrX	147389831	147889899	FMR2(AFF2)	Xq28	Fragile X	300824					
chr1	221168406	221245958	DISP1	1q41	Fryns 1q41	229850					
chr5	112071117	112209835	APC	5q22	Gardner	175100					
chr16	55456620	55507263	SLC12A3	16q13	Gitelman	263800					
chr1	40100001	44100000	GLUT1	1p34.2	Glucose transport defect	606777					
chrX	30581461	30658645	GK	Xp21	Glycerol kinase deficiency	307030					
chr7	41967073	42243900	GLI3	7p14	Greig cephalopolysyndactyly	175700					
chrX	153717263	153904192	F8	Xq28	Hemophilia A	306700					
chrX	138440561	138473283	F9	Xq27	Hemophilia B	306900					
chr6	160341410	160352419	IGF2R	6q25	Hepatocellular carcinoma	114550					
chr18	46810581	46865409	MADH4(SMAD4)	18q21	Hereditary hemorrhagic telangiectasia	175050					
chr9	130300001	133500000	ENG	9q34.11	Hereditary hemorrhagic telangiectasia	187300					
chr2	47630263	47710360	MMR	2p16.3-21	Hereditary non-polyposis colon cancer	120435					
chrX	62771573	62922138	ARHGGEF9	Xq11	Heterotaxy						
chrX	153230091	153256123	FLNA	Xq28	Heterotopia, periventricular, X-linked dominant	300049					
chr2	144100001	148700000	RET	2q22.3	Hirschsprung disease plus	142623					
chr10	42892523	42945803	RET	10q11	Hirschsprung disease plus	142623					
chr13	77367817	77447665	EDNRB	13q22	Hirschsprung disease plus	600155					
chr17	10700000	22200000		17p11.2-p12	HNPP	162500					
chr2	45022365	45025945	SIX3	2p21	Holopresencephaly 2	157170					
chr18	3402072	3448406	TGIF1	18p31	Holopresencephaly 4	142946					
chr13	99432320	99437019	ZIC2	13q32	Holopresencephaly 5	609637					
chr21	46432788	46473119	LSS	21q22	Holopresencephaly 1	236100					
chr21	44256634	44350858	TMEM1	21q22	Holopresencephaly 1	236100					
chr7	155288319	155297728	SHH	7q36	Holopresencephaly 3	142945					
chr9	97245085	97319068	PTCH1	9q22	Holopresencephaly 7	610828					
chr2	121210293	121466899	GLI2	2q14	Holopresencephaly 9	610829					
chr12	171221161	171366482	FBXW11	5q35	Holopresencephaly and preaxial polydactyly	264480					
chr12	113278118	113330630	TBX5	12q14	Holt-Oram	142900					
chrX	152780581	152804778	LICAM	Xq28	HSAS, MASA, CRASH	307000, 303350					
chrX	148368203	148423359	IDS	Xq28	Hunter, Mucopolysaccharidosis type 2	309900					
chrX	172349126	172458979	SLC25A12	2q31	Hypomyelination, global, cerebral	612949					
chrX	21960842	22176399	PHEX	Xp22	Hypophosphatic rickets	307800					
chr15	97010284	97325282	IGF1R	15q25	IGF-1 resistance	147370					
chrX	153423653	153446455	IKBK	Xq28	Incontinentia pigmenti	308300					
chr11	17371008	17455025	ABCC8	11p15	Infantile hypernalinism, enteropathy and deafness (Ushers Disease)	606528 / 276904					
chr11	17472018	17522539	USH1C	11p15	Infantile hypernalinism, enteropathy and deafness (Ushers Disease)	606528 / 276904					
chr15	41710064	41747192	CATSPER2	15q15	Infertility and deafness	611102					
chr15	41612952	41664382	KIAA0377(HISPPD2A)	15q15	Infertility and deafness	611102					
chr6	13646817	135860576	AHI1	6q23	Joubert 3	608692					
chr12	86966921	87060124	CEP290	12q21	Joubert 5	610188					
chr8	11737443	11763055	CTSB	8p23	Kabuki	147920					
chrX	8456915	8660227	KAL1	Xp22.31	Kallmann 1	308700					
chr8	38387813	38445509	FGFR1	8p12	Kallmann 2	147950					
chr8	118890787	119193239	EXT1	8q24.11	Langer-Giedion	150230					
chrX	133421923	133462362	HPRT1	Xq26.2	Lesch-Nyhan	300322					
chr11	67130983	67136581	NDUFV1	11q13	Leukodystrophy						
chr17	7505822	7531642	TP53	17p13.1	Li-Fraumeni 1	151623					
chr7	10289473	103417198	RELN	7q22.1	Lissencephaly with cerebellar hypoplasia	257320					
chr1	156084461	156109878	LMNA	1q22	LMNA	150330					
chr3	30622998	30710637	TGFBR2	3p24	Loeys-Dietz	610380					
chr9	100907233	100956294	TGFBR1	9q22	Loeys-Dietz	609192					
chr11	2412796	2836915	KCNQ1	11p15	Long QT	192500					

chr15	46487797	46725210	FBN1	15q21.1	Marfan 1	154700	•	•	•	•	#
chr3	30647984	3075631	TGFBR2	3p24.1	Marfan 2	154700	•	•	•	•	#
chr22	28329565	28424583	NF2	22q12	Meningioma / NF2	607173/101000	•	•	•	•	#
chrX	77042046	77200000	ATP7A	Xq21.1	Menkes	309400	•	•	•	•	#
chr1	233597351	233678903	TBCE	1q42	Mental retardation		•	•	•	•	#
chr3	46594217	46598956	TDFG1	3p21	Mental retardation		•	•	•	•	#
chr7	142136904	142140495	PRSS1	7q34	Mental retardation		•	•	•	•	#
chr8	1426975	1654048	DLGAP2	8p23	Mental retardation		○	•	•	•	#
chr10	89613175	89718512	PTEN	10q23	Mental retardation		•	•	•	•	#
chr15	23123488	23245220	UBE3A	15q11	Mental retardation		•	•	•	•	#
chr16	971809	976980		16p13	Mental retardation		•	•	•	•	#
chr6	101953675	102623474	GRIK2	6q16	Mental retardation, autosomal recessive 6	611092	○	•	•	•	#
chr22	49410315	49413453	ARSA	22q13	Metachromatic leukodystrophy	250100	•	•	•	•	#
chrX	9500000	17100000	HCSC	Xp22.2	Microphthalmia with linear skin defects	309801	•	•	•	•	#
chr3	182903176	182926972	SOX2	3q26.3-q27	Microphthalmia	206900	•	•	•	•	#
chr17	853509	1037366	ABR	17p13.3	Miller-Dieker	247200	•	•	•	•	#
chr17	2443886	2535638	PAFAH1B1(LUS1)	17p13.3	Miller-Dieker	247200	•	•	•	•	#
chr17	1368033	1412860	PITPNA	17p13.3	Miller-Dieker	247200	•	•	•	•	#
chr17	1194593	1250267	YWHAE	17p13.3	Miller-Dieker	247200	•	•	•	•	#
chr1	68274232	68299047	DIRAS3	1p31	Monosomy 1p31		•	•	•	•	#
chr2	144862055	144994386	ZEB2	2q22	Mowat-Wilson	235730	○	•	•	•	#
chr4	1795039	1810599	FGFR3	4p16.3	Muenke / achondroplasia	602849, 100800	•	•	•	•	#
chr1	46654354	46664121	POMGNT1	1p34.1	Muscle-eye-brain disease	253280	•	•	•	•	#
chrX	149487727	149592272	MTM1	Xq28	Myotubular myopathy	310400	•	•	•	•	#
chr9	128416569	128503132	LXMB1	9q33.3	Nail-patella	161200	•	•	•	•	#
chrX	17303464	17664034	NHS	Xp22	Nance-horan	302350	•	•	•	•	#
chr10	21110094	21503122	NEBL	10p12	Nebulette	605491	•	•	•	•	#
chrX	49573965	49750632	CLCN5	Xp11	Nephrolithiasis 2, X-linked, dent disease	310468	•	•	•	•	#
chr2	110237195	110319883	NPHP1	2q13	Nephronophthisis 1	256100	•	•	•	•	#
chr17	27288185	27352162	JJAZ1(SUZ12)	17q11.2	Neurofibromatosis Type 1	162200	•	•	•	•	#
chr17	26446121	26728821	NF1	17q11.2	Neurofibromatosis Type 1	162200	○	•	•	•	#
chr17	25549032	25586831	SLC6A4	17p11	Neuroticism	607834	•	•	•	•	#
chr1	61320883	61694624	NFIA	1p31.3	NFIA haploinsufficiency	600727	○	•	•	•	#
chr2	111340919	111432099	PTPN11	12q24	Noonan 1	163950	•	•	•	•	#
chr2	3905469	3921067	SOS1	2p22.1	Noonan 4	607333	•	•	•	•	#
chr3	12600108	12690678	RAF1	3p25	Noonan 5	611553	•	•	•	•	#
chrX	43692968	43717694	NDP	Xp11	Norrie	310600	•	•	•	•	#
chr6	100943471	101018272	SIM1	6q16	Obesity, severe		•	•	•	•	#
chr15	25673628	26018061	OCA2	15q13	Oculocutaneous albinism Type 2	203200	○	•	•	•	#
chr20	49833990	49852421	SALL4	20q13	Okhиро	607323	•	•	•	•	#
chr14	36196533	36216762	PAX9	14q13	Oligodontia	604625	•	•	•	•	#
chrX	10376166	10765667	MID1	Xq22	Opitz	300000	•	•	•	•	#
chrX	106956116	107057077	MID2	Xq22	Opitz / FP	300204	•	•	•	•	#
chrX	70255131	70279029	MED12	Xq13.1	Opitz-Kaveggia	305450	•	•	•	•	#
chr3	190831910	191097756	TP73L	3q28	Optic atrophy	165500	•	•	•	•	#
chrX	38096302	38165553	OTC	Xp11.4	Ornithine transcarbamylase deficiency	311250	•	•	•	•	#
chrX	13662801	13697391	OFD1	Xp22.2	Orofaciodigital 1	311200	•	•	•	•	#
chr17	45616456	45633992	COL1A1	17q21	Osteogenesis imperfecta		•	•	•	•	#
chr12	63849638	63928374	LEMD3	12q14.3	Osteopetrolia	166700	•	•	•	•	#
chr12	21300001	26500000		12p12.1	Pallister-killian	601803	•	•	•	•	#
chrX	102918410	102934201	PLP1	Xq22.2	Pelizaeus-merzbacher disease	312080	•	•	•	•	#
chr13	31774112	31906409	B3GALT1	13p13-q12.3	Peters plus	261540	•	•	•	•	#
chr19	1000000	6900000	STK11	19p13.3	Peutz-Jeghers	175200	•	•	•	•	#
chrX	77246425	77268980	PGK1	Xp21	Phosphoglycerate kKinase deficiency	300653	•	•	•	•	#
chr18	51045967	51454183	TCF4	18q21.2	Pitt-Hopkins	610954	○	•	•	•	#
chr4	89147844	89217953	PKD2	4q22.1	Polycystic kidney disease	613095	•	•	•	•	#
chr16	2078712	2125900	PKD1	16p13	Polycystic kidney disease	601313	•	•	•	•	#
chr7	15168394	156676602	LMBR1	7q36	Polycystic preaxial 2	174500	•	•	•	•	#
chr17	25800000	31800000		17q11.2	Potocki-Lupski	610883	•	•	•	•	#
chr11	44242734	44288292	ALX4	11p11.2	Potocki-Shaffer	601224	•	•	•	•	#
chr11	44073675	44223555	EXT2	11p11.2	Potocki-Shaffer	601224	○	•	•	•	#
chrX	1	2700000	PAR1	Xp22	Pseudoautosomal		•	•	•	•	#
chrX	154595003	154913754	PAR2	Xq28	Pseudoautosomal		•	•	•	•	#
chr2	31603160	31695944	SRD5A2	2p23	Pseudovaginal perineoscrotal dysoypadias	264600	•	•	•	•	#
chrX	19271972	19287886	PDHA1	Xp22	Pyruvate dehydrogenase deficiency	312170	•	•	•	•	#
chr17	33120548	33179182	TCF2(HNF1B)	17q12	Renal cysts and diabetes	137920	•	•	•	•	#
chr13	47775912	47954023	RB1	13q14.2	Retinoblastoma	180200	•	•	•	•	#
chrX	152940458	153016323	MECP2	Xq28	Retts	312750	•	•	•	•	#
chr4	111758029	111782566	PITX2	4q25	Rieger	180500	•	•	•	•	#
chrX	99785819	99812952	SRPX2	Xq22	Rolandic epilepsy, X-linked mental retardation, and speech dyspraxia	300643	•	•	•	•	#
chr16	3717720	3870723	CREBBP	16p13	Rubinstein taybi	610543	•	•	•	•	#
chr16	3642941	3648097	DNAH1	16p13	Rubinstein taybi	610543	•	•	•	•	#
chr7	19121616	19129550	TWIST1	7p21.1	Saethre-Chozen	101400	•	•	•	•	#
chr10	119291946	119299043	EMX2	10q26.11	Schneppcephaly	289160	•	•	•	•	#
chr17	39186625	39191682	SOST	17q21.31	Sclerostin	269500	•	•	•	•	#
chr2	166553916	166638395	SCN1A	2q24.3	SCN1A related seizures	609634	•	•	•	•	#
chr1	178466065	178510811	LHX4	1q25.2	Short stature	282700	•	•	•	•	#
chrX	505079	540146	SHOX	Xp22.33	SHOX	312865	•	•	•	•	#
chrX	50615258	50838651	GRB10	7p12	Silver-Russell	180860	•	•	•	•	#
chrX	132497442	132947332	GPC3	Xq26	Simpson, golabi, behmel, type 1	312870	○	•	•	•	#
chr11	70823105	70837125	DHCR7	11q13.4	Smith-leml-opitz	270400	•	•	•	•	#
chr17	17525512	17655488	RAI1	17p11.2	Smith-magenis	182290	•	•	•	•	#
chr5	176492686	176659820	NSD1	5q35.3	Sotos	117550	•	•	•	•	#
chr7	96477643	96502078	DLX5	7q21.3	Split hand and foot	220600	•	•	•	•	#
chr7	96140173	96177139	SHFM1	7q21.3	Split hand and foot	220600	•	•	•	•	#
chr10	103103815	103307060	BTRC	10q24.32	Split hand and foot malformation 3	600095	•	•	•	•	#
chr10	103360412	103444733	FBXW4	10q24.32	Split hand and foot malformation 3	600095	•	•	•	•	#
chr10	102976723	102978707	LBX1	10q24.32	Split hand and foot malformation 3	600095	•	•	•	•	#
chr10	103328629	103338004	PDLL	10q24.32	Split hand and foot malformation 3	600095	•	•	•	•	#
chr2	172658454	172662647	DLX1	2q31	Split hand and foot malformation 5	606708	•	•	•	•	#
chr2	172672412	172675724	DLX2	2q31	Split hand and foot malformation 5	606708	•	•	•	•	#
chrX	7075293	7282680	STS	Xp22.31	Steroid sulfatase	308100	•	•	•	•	#
chrX	6461659	6463159	VXCA3A	Xp22.31	Steroid sulfatase	308100	•	•	•	•	#
chr12	46653018	46684528	COL2A1	12q13.11	Stickler 1	108300	•	•	•	•	#
chr1	103115583	103346640	COL11A1	1p21	Stickler 2	604841	•	•	•	•	#
chr2	176653081	176656936	EVX2	2q31	Synpolydactyly	186000	•	•	•	•	#
chr2	17665778	176668912	HOXD13	2q31.1	Synpolydactyly	186000	•	•	•	•	#
chr2	176655334	176697891	HOXD9	2q31	Synpolydactyly	186000	•	•	•	•	#
chr2	176664382	176767892	HOXDgenes	2q31	Synpolydactyly	186000	•	•	•	•	#
chr17	42686207	42745076	ITGB3	17q21	Thrombosthenia of glanzmann and naegeli	273800	•	•	•	•	#
chr1	145841561	145848017	GJA8	1q21.1	Thrombocytopenia absent radius	274000	•	•	•	•	#
chr12	1950490	2677376	CACNA1C	12p13	Timothy	601005	○	•	•	•	#
chr16	49727387	49742684	SALL1	16q12.1	Townes brooks	107480	•	•	•	•	#
chr8	116489900	116750429	TRPS1	8q23.3	Trichorhino-inophlangeal, langer-giedion	150230	•	•	•	•	#
chr9	134756557	134809841	TSC1	9q34.13	Tuberous sclerosis 1	181100	•	•	•	•	#
chr16	2038600	2078713	TSC2	16p13.3	Tuberous sclerosis 2	613254	•	•	•	•	#
chr12	113592442	113606362	TBX3	12q24.21	Ulnar-mammary	181450	•	•	•	•	#
chr1	20027895	200404102	IRF6	1q32.2	Van der Woude	119300	•	•	•	•	#
chr6	43845931	43862201	VEGFA	6p21	Vascular endothelial growth factor disorders		•	•	•	•	#
chr7	113513618	114118328	FOXP2	7q31.1	Verbal dyspraxia	602081	•	•	•	•	#
chr2	130995137	131002053	CFC1	2q21	Visceral heterotaxy	605376	•	•	•	•	#
chr3	10158319	10168746	VHL	3p25.3	Von hippel lindau	193300	•	•	•	•	#
chr22	38368319	38380539	SOX10	22q13.1	Wardenberg 4	609136	•	•	•	•	#
chr2	222772851	222871944	PAX3	2q36.1	Wardenberg I	193500	•	•	•	•	#
chr3	69871323	70100177	MITF	3p14.1	Wardenberg IIA	193510	•	•	•	•	#
chr11	32355900	32423662	WT1	11p13	WAGR	194072	•</				

chrX	72863307	73063491	XIST	Xq13	X inactivation specific transcript		●	●	●	
chrX	153644344	153659154	DKC1	Xq28	X-linked dyskeratosis congenita	305000	●	●	●	#
chrX	136476012	136481923	ZIC3	Xq26.3	X-linked heterotaxy	306955	●	●	●	#
chrX	18567731	18600150	RS1	Xp22	X-linked juvenile retinoschisis	312700	●	●	●	#
chrX	123307875	123334696	SH2D1A	Xq25	X-linked lymphoproliferative type 1	308240	●	●	●	#
chrX	122821729	122875503	BIRC4(XIAP)	Xq25	X-linked lymphoproliferative type 2	308240	●	●	●	#
chrX	108771220	108863275	ACSL4	Xp22	X-linked mental retardation		●	●	●	
chrX	115216031	115219848	AGTR2	Xq23	X-linked mental retardation		●	●	●	
chrX	15753850	15783021	AP1S2	Xp22	X-linked mental retardation		●	●	●	
chrX	135575377	135691169	ARHGEF6	Xq26	X-linked mental retardation		●	●	●	
chrX	24932213	24943775	ARX	Xp21	X-linked mental retardation		●	●	●	
chrX	40325160	40350832	ATP6AP2	Xp11	X-linked mental retardation		●	●	●	
chrX	39795561	39921526	BCOR	Xp11	X-linked mental retardation		●	●	●	
chrX	79818351	79951889	BRWD3	Xp21	X-linked mental retardation		●	●	●	
chrX	41264287	41667212	CASK	Xp11	X-linked mental retardation		●	●	●	
chrX	119542474	119593712	CUL4B	Xq24	X-linked mental retardation		●	●	●	
chrX	69581449	69642062	DLG3	Xq13	X-linked mental retardation		●	●	●	
chrX	47379864	47394994	ELK1	Xp11	X-linked mental retardation		●	●	●	
chrX	14771450	14801105	FANCB	Xp22	X-linked mental retardation		●	●	●	
chrX	48219493	48229696	FTSJ1	Xp11	X-linked mental retardation		●	●	●	
chrX	153318715	153325009	GDI1	Xq28	X-linked mental retardation		●	●	●	
chrX	122145839	122450474	GRIA3	Xq24	X-linked mental retardation		○	●	●	
chrX	53474931	53478048	HADH2(HSD17B10)	Xp11	X-linked mental retardation		●	●	●	
chrX	11039373	11051122	HCCS	Xp22	X-linked mental retardation		●	●	●	
chrX	28515480	29884757	IL1RAPL1	Xp21	X-linked mental retardation		●	●	●	
chrX	53237378	53271329	JARID1C	Xp11	X-linked mental retardation		●	●	●	
chrX	73870137	74061709	KIAA2022	Xq13	X-linked mental retardation		○	●	●	
chrX	56275632	56328254	KLF8	Xp11	X-linked mental retardation		●	●	●	
chrX	118889762	118894646	NDUFA1	Xq24	X-linked mental retardation		●	●	●	
chrX	100973741	100999205	NXF5	Xq22	X-linked mental retardation		●	●	●	
chrX	110074169	110350829	PAK3	Xp22	X-linked mental retardation		○	●	●	
chrX	53979838	54088121	PHF8	Xp11	X-linked mental retardation		●	●	●	
chrX	48252315	48264146	PORCN	Xp11	X-linked mental retardation		●	●	●	
chrX	48640139	48645364	PQBP1	Xp11	X-linked mental retardation		●	●	●	
chrX	153279912	153283874	RPL10	Xq28	X-linked mental retardation		●	●	●	
chrX	50351387	50573784	SHROOM4(KIAA1202)	Xp11	X-linked mental retardation		●	●	●	
chrX	48201871	48213509	SLC38A5	Xp11	X-linked mental retardation		●	●	●	
chrX	21868763	21922876	SMS	Xp22	X-linked mental retardation		●	●	●	
chrX	139411973	139415436	SOX3	Xq27	X-linked mental retardation		●	●	●	
chrX	47316244	47364200	SYN1	Xp11	X-linked mental retardation		●	●	●	
chrX	38305683	38433116	TM4SF2(TSPAN7)	Xp11	X-linked mental retardation		●	●	●	
chrX	118592527	118602407	UBE2A	Xq23	X-linked mental retardation		●	●	●	
chrX	118852017	118870996	UPF3B	Xq24	X-linked mental retardation		○	●	●	
chrX	8393346	8394346	VCX3A	Xp22	X-linked mental retardation		●	●	●	
chrX	74508786	74659600	ZDHC15	Xq13	X-linked mental retardation		●	●	●	
chrX	128766594	128805805	ZDHC9	Xq25	X-linked mental retardation		●	●	●	
chrX	70376199	70391660	ZNF261(ZMYM3)	Xq13	X-linked mental retardation		●	●	●	
chrX	47191350	47227289	ZNF41	Xp11	X-linked mental retardation		●	●	●	
chrX	47581245	47666550	ZNF81	Xp11	X-linked mental retardation		●	●	●	
chrX	73557810	73670475	SLC16A2	Xq13	X-linked mental retardation, allan-herndon-dudley	300523	●	●	●	#
chrX	134895252	134957094	SLC9A6	Xq26	X-linked mental retardation, christianson type	300243	●	●	●	#
chrX	53576476	53793398	HUWE1	Xp11	X-linked mental retardation, turner	300706	○	●	●	#
chrX	67179440	67570372	OPHN1	Xq12	X-linked mental retardation, with cerebellar hypoplasia and distinctive facial appearance	300486	●	●	●	#
chrX	18353678	18581666	CDKL5	Xp22.13	X-linked spasms	300672	●	●	●	#
chrX	46581319	46626733	RP2	Xp11	Xp11.3 deletion	300578	●	●	●	#
chrX	46243490	46289820	ZNF674	Xp11	Xp11.3 deletion	300578	●	●	●	#
chr9	126283337	126309520	NR5A1	9q33.3	XY Sex Reversal	184757	●	●	●	#
chr19	62384680	62432350	ZNF264	19q13	Zinc finger protein-264		●	●	●	
chr2	202949916	203140719	BMPR2	2q32			●	●	●	
chr7	129710229	129761249	CPA4	7q32			●	●	●	
chr7	130058017	130079399	KLF14	7q32			●	●	●	
chr7	129903281	129943368	MEST	7q32			●	●	●	
chr7	98084545	98097116	NPTX2	7q22			●	●	●	
chr7	94364911	94770958	PPP1R9A	7q21			○	●	●	
chr8	140683985	140794480	KCNK9	8q24			●	●	●	
chr11	1975651	1980951	H19_ICRegion	11p15			●	●	●	
chr11	2671987	2683633	KCNQ1_ICRegion	11p15			○	●	●	
chr11	3054921	3153115	OSBPL5	11p15			●	●	●	
chr13	65774967	66702464	PCDH9	13q21			●	●	●	
chr14	20923198	20975244	CHD8	14q11			●	●	●	
chr14	100252981	100281225	DLK1	14q32			●	●	●	
chr14	100261144	100364369	DLK1&MEG3_ICRegion	14q32			●	●	●	
chr14	28306038	28308622	FOXP1B	14q12			●	●	●	
chr14	100352214	100407118	MEG3	14q32			●	●	●	
chr14	20897000	20932000	MicroDeletionRegion	14q11			●	●	●	
chr14	20889476	20922265	SUPT16H	14q11			●	●	●	
chr15	49527231	49702259	DMXL2	15q21			●	●	●	
chr16	6009133	7702500	A2BP1	16p13			●	●	●	
chr18	73090996	73111084	GALR1	18q23			●	●	●	
chr18	42798570	42820446	TCEB3C	18q21			●	●	●	
chr19	62005614	62053875	PEG3	19q13			●	●	●	
chr19	62040958	62043979	PEG3_ICRegion	19q13			●	●	●	
chr19	61967731	62053886	ZIM2	19q13			●	●	●	
chr20	56876787	56880787	GNAS_ICRegion	20q13			●	●	●	
chr20	56859055	56861251	GNAS_ICRegion	20q13			●	●	●	
chr20	56896896	56899356	GNAS_ICRegion	20q13			●	●	●	
chr20	41566487	41613948	L3MBTL	20q13			●	●	●	
chr20	35573020	35595505	NNAT	20q11			●	●	●	
chr22	16454903	16491588	ATP6V1E1	22q11			●	●	●	
chrY	2714821	2716600	SRY	Yp11.31			●	●	●	#

Microdeletions or microduplications have been associated with this condition

○ Low coverage of syndrome

● Coverage of syndrome

○ No Coverage of Syndrome

Each design includes coverage for all aneuploidies and all sub-telemeres

Summary for each array, plus coverage within each gene/targeted region:

Application	No. of arrays/slide	No. of probes	Genome/backbone probe density	No. of targeted regions	Probe resolution	Scanner compatibility	Product name
Whole genome screening	8	60,000	Probe every 60kb	498	Probe every 31kb within genes, 4 probes per gene (on average)	High res.	CytoSure ISCA v2 (8x60k)
Whole genome screening	4	180,000	Probe every 25kb	501	Probe every 14kb within genes, 6 probes per gene (on average)	High res.	CytoSure ISCA v2 (4x180k)
Whole genome screening	2	105,000	Probe every 40kb	610	Probe every 3kb within genes, 33 probes per gene (on average)	High res. & 5 μm	CytoSure Syndrome Plus v2 (2x105k)
X chromosome analysis	4	44,000	Probe every 4kb	1118	Probe every 380bp within genes, 81 probes per gene (on average)	High res. & 5 μm	CytoSure Chromosome X (4x44k)

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