



CPT CODES REQUIRING PRIOR AUTHORIZATION

Beginning November 1, 2017, UnitedHealthcare (UPC) has announced that the following CPT codes will require PRIOR AUTHORIZATION before submitting the orders to the performing laboratory.

Metro Lab Ordering Unit Code	Test Name	CPT Code
10149	Cystic Fibrosis 165 Pathog.Variants	81220
50045	FACTOR V LEIDEN (R506Q) MUTATION	81241
50055	PROTHROMBIN G 20210A GENE MUTATION	81240
157511	FLT3 & NPM1 MUTATION DETECTION	81245
181403	Bill: RhD Variet Assay	81403
181479	Bill: RBC Molecular Phenotype	81479
226022	BILL: INMS Unlisted Molec.Path	81479
281240	Billing: F2 Gene	81240
281310	BILL: NPM1 GENE ANALY EXON 12 VAR	81310
290467	Y Chromosome Microdeletions/PCR	81403
292032	CYP2D6 Genotype	81226
292035	JAK2 V617F Mutation Detect. Rpt:	81270
292331	CYP2D6 Genotype - Tamoxifen	81226
297102	SPINAL MUSC.ATROPHY CARRIER TEST	81401
300184	Twin Zygosity Testing	81265
300550	Rflx: Factor V Leiden(F5) R506Q	81241
300566	Prothrombin (F2) G20210A Mut.Rflx	81240
302481	Hemochromatosis (HFE) 3 Mutation	81256
302696	Celiac (HLA-DQ2 & DQ8) Genotyping	81376
302697	Billing: HLA DQB1 allele(s)	81383
302891	HLA-B*57:01,Abicavir Sensitivity	81381
302964	JAK2 V617F QL,Bld/B.Marrow/Rfl PCR	81270
302966	JAK2 V617F QL,Bld/Bone Marrow	81270
303035	Narcolepsy (HLA-DQB1*06:02)Genotype	81383
303341	HLA Class I (ABC) Sequencing	81379
303441	BCR/ABL1, Major (p210),Quantitative	81206
303651	HLA-DQB Genotyping	81382
303861	Alpha Thalassemia, 7 Deletions	81257
303891	Huntington Disease (HD) by PCR	81401
303892	Billing: Target Seq Molec Path Proc	81479
303921	Hemophilia A (F8) 2 Inversions	81403
303926	Hemophilia A (F8) Comprehen.Rpt.	81403
304021	bcr/abl,t(9;22)Qual.Transl.RT-PCR	81206
304050	FLT3 Mutation Detection by PCR	81245
304055	NPM1 Mutation Detection by PCR	81310
304161	IL28B-Associated Variants,2 SNPs	81400
304301	Ankylosing Spondylitis Rpt:	81374

Metro Lab Ordering Unit Code	Test Name	CPT Code
304351	MEN2 (RET) Gene Mutat./Seq,Bld	81405
304376	Marfan Syndrome (FBN1) Sequencing	81408
304420	Cytogenomic SNP(aCGH) Microarray	81229
304580	MCAD Deficiency(ACADM)Sequencing,P	81479
304585	MCAD(ACADM) 2 Mutations (PCR),P	81401
305041	BCR-ABL1 Mutation Analysis/NGS	81479
305146	Bill: Molec.Path.Proc DNA seq/pcr	81403
305355	Y Chromosome Microdeletion	81403
305455	Fragile X (FMR1) w/Rflx Methylation	81244
305600	Prenatal Testing Fetal Aneuploidy	81420
305605	Spinobulbar Musc Atrophy,Kennedy's	81401
305770	Plasminogen Activ.Inhib-1,Genotype	81400
305900	bcr/abl,t(9;22)Qual.Transl/Rfl QN	81206
306016	MTHFR Mutation Detection-2 Mutation	81291
306150	Ashkenazi Jewish Pnl 16 Gene Rpt:	81200
306190	HLA-B*1502 Typing	81381
306260	HLA-B51 Typing	81381
306265	Ehlers-Danlos Type IV (COL3A1 Gene)	81479
306345	Ehlers-Danlos Syndr IV (COL3A1):	81408
306365	CYP450 2D6(CYP2D6) 15var, Gene Dup	81226
306400	A1A Genotype /w Rflx Phenotype	81332
306510	Platelet AG 1 Genotyping (HPA-1)	81400
306745	FAP:APC,Seq,Del/Dup(MUTYH) 2 Mut	81201
306750	Connexin 26 (GJB2),Sequencing,Bld	81252
306875	T-Cell Clonality Screening by PCR	81342
306990	CALR Exon 9 Mutation Analysis/PCR	81219
307005	Aortopathy Panel Rpt:	81410
307030	JAK2 Exon 12 Mutation Analysis/PCR	81403
307095	Renal Hereditary Cancer Pnl Rpt:	81445
307115	Pancreatitis,Pnl, Gene Seq. Rpt:	81223
307180	Angelman/Prader-Willi Syndr/M-S PCR	81331
307240	IGHV Mutation Analysis/Sequencing	81263
307270	TPMT Genotype (Prometheus)	81401
307275	5-FU Mutations,Toxicity/Response	81400
307345	Neurofibromatosis 1 Seq,DelDup	81408
307350	HMBS Gene,Full Gene Analysis	81479
307425	HLA-B Genotype	81380

Metro Lab Ordering Unit Code	Test Name	CPT Code
307560	PML-RARA Transloc.t(15:17),PCR/QNT	81315
307615	Neurofibromatosis NF2 Seq,Del/Dup	81405
307620	MPL Exon 10 Mutation Detection,Bld	81403
307625	CYP21A2 Full Gene Analysis,Bld	81405
307630	JAK2 V617F QNT,Bld	81270
307690	Cytogenomic SNP Microarray-Oncology	81406
307700	Breast & Ovarian(BRCA1/BRCA2)FGA	81162
307710	Cystic Fibros.165 Var. w/Rflx Seq.	81220
307715	MYD88 L265P Mutation Detect.QN/PCR	81479
307730	CF (CFTR)Seq/Rflx Del/Dup(2nd Tier)	81223
307755	MPL codon 515 Mut.Detect/Pyroseq.QN	81402
307765	COL4A1 DNA Sequencing (CSVD),Bld	81408
307855	Factor V,R2 Mutation by PCR	81400
308025	Von Hippel-Lindau(VHL) Gene Analy.	81404
308055	EGFR T790M Mutation Detection/PCR	81235
308065	Kit Mutations in AML,Frag.Analy/Seq	81272
308070	CEBPA Mutation Detection	81218
308100	Beta Globin (HBB) Gene Sequencing	81404
308110	UGT1A1 Genotyping	81350
308155	Epi proColon	81327
308175	Charcot-Marie-Tooth(1A)HNPP Del/Dup	81324
308200	Marfan Syndrome(FBN1)Seq &Del/Dup	81479
308220	HLA-DP Genotyping	81382
308225	HLA-DQ Genotyping	81382
308235	TPMT Genotyping,4 Variants,Blood	81401
308265	DPYD,3 Varients,Genotype,Blood	81400
308310	Cardiomyopathy/Arrhythm.Seq.85gene	81413
308320	HLA-DRB1 by Next Generation Seq.	81382
308325	HLA DRB 3-,4-,5-	81382
308390	TP53 Somatic Mutation,Prognostic	81405
308445	MEN Type 1 Sequencing	81405
308465	Custom Sequencing by Consultation	81479
371404	BILL: ALPHA GLOB(HBA1/HBA2)DEL/DUP	81404
371405	BILL: Alpha Thallass.HBA1/HBA2 Seq	81405
371479	BILL: HGB LEPORE HBD/HBB 3 MUTAT.	81479
380403	BILL: BETA GLOB(HBB) DEL/DUP	81403
381202	Rflx: APC Targeted Sequencing Bill	81202
381203	BILL: APC GENE DUPL/DEL VARIENTS	81203
381207	Billing: BCR/ABL1 Minor Brkpt	81207
381209	Billing: Bloom Gene Analysis	81209
381213	BILL: BRCA1&2 UNCOMM DUP DEL VAR	81213
381215	BILL: BRCA1 Target Seq Gene	81215
381217	BILL: BRCA2 Target Seq Gene	81217
381221	Rflx:CFTR Targeted Sequencing Bill	81221
381222	BILL: CFTR Deletion/Duplication	81222
381223	BILL: Molec.Path.Full Gene Seq.	81223
381240	BILLING F2 GENE ANALYS 20210G-A	81240

Metro Lab Ordering Unit Code	Test Name	CPT Code
381241	BILLING F5 GENE ANALY LEIDEN VAR	81241
381242	Billing: FANCC Gene Analysis	81242
381244	BILL: FMR1 Methylation by mPCR	81244
381246	BILL: TKD Varients	81246
381251	Billing: GBA Gene Analysis	81251
381253	Rflx:GJB2 Targeted Sequencing Bill	81253
381255	Billing: HEXA Gene Analysis	81255
381260	Billing: IKBKAP Gene Analysis	81260
381290	Billing: MCOLN1 Gene Analysis	81290
381292	BILL: MLH1 GENE ANALYS,FULL SEQ	81292
381293	Rflx:MLH1 Targeted Sequencing Bill	81293
381294	BILL: MLH1 GENE ANALY DUP/DEL VAR	81294
381295	BILL: MSH2 FULL GENE/FULL SEQ	81295
381296	Rflx:MSH2 Targeted Sequencing Bill	81296
381297	BILL: MSH2 GENE ANALY,DUP/DEL VAR	81297
381298	BILL: MSH6 GENE ANALY,FULL SEQ	81298
381299	Rflx:MSH6 Targeted Sequencing Bill	81299
381300	BILL: MSH6 GENE ANALY,DUP/DELET VAR	81300
381303	Rflx:MECP2 Targeted Sequencing Bill	81303
381318	Rflx:PMS2 Targeted Sequencing Bill	81318
381322	Rflx:PTEN Targeted Sequencing Bill	81322
381323	BILL: PTEN GENE ANALY,DUP/DEL VAR	81323
381330	Billing: SMPD1 Gene Analysis	81330
381401	Bill: Molecular Pathology,Level2	81401
381402	BILL: MOPATH Procedure Level 3	81402
381403	BILL: Molec.Path Proced.LV 4	81403
381404	BILL: Molecular Path.Proc Level 5	81404
381405	Bill: Molec.Path Level 6	81405
381406	Bill: Molec.Path Level 7	81406
381407	BILL: MOLEC PATH PROCEDURE LEVEL 8	81407
381408	Bill: Molec.Path Level 9	81408
381411	BILL: AORTIC DYSF/DILAT 4 GENE	81411
381479	BILL: Unlisted MolecPath Proced.	81479
382205	BILL: BCKDHB GENE ANALYSIS	81205
382221	RFLX:CFTR TARGETED SEQUEN	81221
382250	BILL: G6PC GENE ANALYSIS	81250
382400	BILL: MOLECULAR PATH LEVEL I	81400
382401	BILL: BILL: MOLECULAR PATH LEVEL II	81404
382479	BILL: UNLISTED MOLEC.PATH	81479
391401	BILL: Beta Globulin Mutations	81401
391402	BILL: MEFV81404 Target Seq Gene	81402
391403	BILL: HEREDIT.PERSIST.FETAL HGB	81403
391404	BILL: MOLEC PATH PROCEDURE LEVEL 5	81404
391405	Bill: Molec.Path.Proced.Level 6	81405
391406	BILL: MOPATH Level 7	81406
391408	Bill: Molec.Path Level 9	81408
391479	BILL: Unlisted Molec.Path.	81479

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