Total Approvals 253

This report will be updated January and July of every year

NDA and BLA Accelerated Approvals

Application Number	Proprietary Name	Established Name	Applicant	FDA Received Date	Accelerated Approval Date	Total Time to Accelerated Approval (Months)	Accelerated Approval Indication	Conversion-Withdrawal Status	Full Approval Conversion- Withdrawal Da
NDA 214701	GAVRETO	PRALSETINIB	BLUEPRINT MEDICINES CORP	6/30/2020	12/1/2020	5.1	FOR THE TREAMENT OF ADULT AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER WITH ADVANCED OR METASTATIC RET-MUTANT MEDULLARY THYROID CANCER (MTC) WHO REQUIRE SYSTEMIC THERAPY	Not Yet Converted	
NDA 214701	GAVRETO	PRALSETINIB	BLUEPRINT MEDICINES CORP	6/30/2020	12/1/2020	5.1	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER WITH ADVANCED OR METASTATIC RET FUSION-POSITIVE THYROID CANCER WHO REQUIRE SYSTEMIC THERAPY AND WHO ARE RADIOACTIVE IODINE-REFRACTORY (IF RADIOACTIVE IODINE IS APPROPRIATE)		
BLA 761171	DANYELZA	NAXITAMAB-GQGK	Y-MABS THERAPEUTICS INC	3/31/2020	11/25/2020	7.9	IN COMBINATION WITH GRANULOCYTE-MACROPHAGE COLONY-STIMULATING FACTOR (GM-CSF), FOR THE TREATMENT OF PEDIATRIC PATIENTS 1 YEAR OF AGE AND OLDER AND ADULT PATIENTS WITH RELAPSED OR REFRACTORY HIGH-RISK NEUROBLASTOMA IN THE BONE OR BONE MARROW WHO HAVE DEMONSTRATED A PARTIAL RESPONSE, MINOR RESPONSE, OR STABLE DISEASE TO PRIOR THERAPY	Not Yet Converted	
BLA 125514 Supplement 88	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME CORP	5/28/2020	11/13/2020	5.6	IN COMBINATION WITH CHEMOTHERAPY, FOR THE TREATMENT OF PATIENTS WITH LOCALLY RECURRENT UNRESECTABLE OR METASTATIC TRIPLE NEGATIVE BREAST CANCER WHOSE TUMORS EXPRESS PD-L1 [COMBINED POSITIVE SCORE (CPS) 2+0] AS DETERMINED BY AN FDA APPROVED TEST	Not Yet Converted	
NDA 213721	GAVRETO	PRALSETINIB	BLUEPRINT MEDICINES CORP	3/23/2020	9/4/2020	5.4	FOR THE TREATMENT OF ADULT PATIENTS WITH METASTATIC RET FUSION-POSITIVE NON-SMALL CELL LUNG CANCER (NSCLC) AS DETECTED BY AN FDA APPROVED TEST	Not Yet Converted	
NDA 212154	VILTEPSO	VILTOLARSEN	NIPPON SHINYAKU CO LTD□	12/12/2019	8/12/2020	8.0	FOR THE TREATMENT OF DUCHENNE MUSCULAR DYSTROPHY (DMD) IN PATIENTS WHO HAVE A CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 53 SKIPPING FOR THE TREATMENT OF CHAGAS DISEASE IN	Not Yet Converted	
NDA 213464	LAMPIT	NIFURTIMOX	BAYER HEALTHCARE PHARMACEUTICALS	12/6/2019	8/6/2020	8.0	PEDIATRIC PATIENTS BIRTH TO LESS THAN 18 YEARS OF AGE AND WEIGHING AT LEAST 2.5 KG	Not Yet Converted	
BLA 761158	BLENREP	BELANTAMAB MAFODOTIN-BLMF	GLAXOSMITHKLINE INTELLECTUAL PROPERTY DEVELOPMENT LTD ENGLAND	12/5/2019	8/5/2020	8.0	FOR THE TREATMENT OF ADULTS WITH RELAPSED OR REFRACTORY MULTIPLE MYELOMA WHO HAVE RECEIVED AT LEAST FOUR PRIOR THERAPIES INCLUDING AN ANTI-CD38 MONOCLONAL ANTIBODY, A PROTEASOME INHIBITOR, AND AN IMMUNOMODULATORY AGENT	Not Yet Converted	
BLA 761163	MONJUVI	TAFASITAMAB- CXIX	MORPHOSYS US INC	12/30/2019	7/31/2020	7.0	IN COMBINATION WITH LENALIDOMIDE FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY DIFFUSE LARGE B-CELL LYMPHOMA (DLBCL) NOT OTHERWISE SPECIFIED, INCLUDING DLBCL ARISING FROM LOW GRADE LYMPHOMA, AND WHO ARE NOT ELIGIBLE FOR AUTOLOGOUS STEM CELL TRANSPLANT (ASCT)	Not Yet Converted	
BLA 125514			MERCK SHARP &				PROVIDES FOR AN ALTERNATE DOSAGE REGIMEN OF 400 MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH RECURRENT OR METASTATIC CUTANEOUS SQUAMOUS CELL CARCINOMA (CSCC) THAT IS NOT CURABLE BY		
Supplement 92	KEYTRUDA	PEMBROLIZUMAB	DOHME	6/12/2020	6/24/2020	0.4	SURGERY OR RADIATION	Not Yet Converted	1

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NDA 212306 Supplement 1	XPOVIO	SELINEXOR	KARYOPHARM THERAPEUTICS INC	12/23/2019	6/22/2020	6.0	FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY DIFFUSE LARGE B-CELL LYMPHOMA (DLBCL), NOT OTHERWISE SPECIFIED, INCLUDING DLBCL ARISING FROM FOLLICULAR LYMPHOMA, AFTER AT LEAST 2 LINES OF SYSTEMIC THERAPY	Not Yet Converted	
NIDA 242400	TA7\/FDII/	TAZEMETOSTAT	EDIZVME INC	42/40/2040	0/40/2020		FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY (R/R) FOLLICULAR LYMPHOMA (FL) WHOSE TUMORS ARE POSITIVE FOR AN EZH2 MUTATION AS DETECTED BY AN FDA-APPROVED TEST AND WHO HAVE RECEIVED AT LEAST 2 PRIOR SYSTEMIC THERAPIES; and THE TREATMENT OF ADULT PATIENTS WITH RIR FL WHO HAVE NO SATISFACTORY AND THE TREATMENT OF THE STATE OF	Net Ver Commented	
NDA 213400 BLA 125514 Supplement 90	TAZVERIK KEYTRUDA	TAZEMETOSTAT PEMBROLIZUMAB	MERCK SHARP & DOHME	12/18/2019	6/18/2020 6/16/2020	6.0	ALTERNATIVE TREATMENT OPTIONS. PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400 MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH UNRESECTABLE OR METASTATIC TMB-H [≥10 MUTATIONS/MEGABASE (MUTMB)] SOLID TUMORS, AS DETERMINED BY AN FDA-APPROVED TEST, THAT HAVE PROGRESSED FOLLOWING PRIOR TREATMENT AND WHO HAVE NO SATISFACTORY ALTERNATIVE TREATMENT OPTIONS	Not Yet Converted Not Yet Converted	
BLA 125514 Supplement 71	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	12/16/2019	6/16/2020	6.0	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS WITH UNRESECTABLE OR METASTATIC TUMOR MUTATIONAL BURDEN-HIGH (TMB-H) [210 MUTATIONS/MEGABASE (MUT/MB)] SOLID TUMORS, AS DETERMINED BY AN FDA-APPROVED TEST, THAT HAVE PROGRESSED FOLLOWING PRIOR TREATMENT AND WHO HAVE NO SATISFACTORY ALTERNATIVE TREATMENT OPTIONS	Not Yet Converted	
NDA 213702	ZEPZELCA	LURBINECTEDIN	JAZZ PHARMACEUTICALS IRELAND LTD	12/16/2019	6/15/2020	6.0	FOR THE TREATMENT OF ADULT PATIENTS WITH METASTATIC SMALL CELL LUNG CANCER (SCLC) WITH DISEASE PROGRESSION ON OR AFTER PRIOR PLATINUM BASED CHEMOTHERAPY	Not Yet Converted	
NDA 204384 Supplement 13	SIRTURO	BEDAQUILINE	JANSSEN RESEARCH AND DEVELOPMENT	11/27/2019	5/27/2020	6.0	FOR THE TREATMENT OF PULMONARY MULTI-DRUG RESISTANT TUBERCULOSIS AS PART OF COMBINATION THERAPY, IN ADULT AND PEDIATRIC PATIENTS (12 TO LESS THAN 18 YEARS OF AGE AND WEIGHING AT LEAST 30 KG) TO INCLUDE PATIENTS ≥ 5 TO <12 YEARS OF AGE AND WEIGHING AT LEAST 15 KG	Not Yet Converted	
NDA 212269	FERRIPROX	DEFERIPRONE	CHIESI USA INC	7/19/2019	5/19/2020	10.0	FOR THE TREATMENT OF PATIENTS WITH TRANSFUSIONAL IRON OVERLOAD DUE TO THALASSEMIA SYNDROMES WHEN CURRENT CHELATION THERAPY IS INADEQUATE	Not Yet Converted	
NDA 209115 Supplement 4	RUBRACA	RUCAPARIB	CLOVIS ONCOLOGY	11/15/2019	5/15/2020	6.0	FOR THE TREATMENT OF ADULT PATIENTS WITH A DELETERIOUS BRCA MUTATION (GERMILINE AND/OR SOMATIC)-ASSOCIATED METASTATIC CASTRATION- RESISTANT PROSTATE CANCER WHO HAVE BEEN TREATED WITH ANDROGEN RECEPTOR-DIRECTED THERAPY AND A TAXANE-BASED CHEMOTHERAPY	Not Yet Converted	
NDA 204026 Supplement 24	POMALYST	POMALIDOMIDE	CELGENE	11/19/2019	5/14/2020	5.8	FOR THE TREATMENT OF KAPOSI'S SARCOMA IN PATIENTS WHO ARE HIV-NEGATIVE	Not Yet Converted	
NDA 204026 Supplement 23	POMALYST	POMALIDOMIDE	CELGENE	11/14/2019	5/14/2020	6.0	FOR THE TREATMENT OF ADULT PATIENTS WITH AIDS- RELATED KAPOSI SARCOMA (KS) AFTER FAILURE OF HIGHLY ACTIVE ANTIRETROVIRAL THERAPY (HAART)	Not Yet Converted	
NDA 213246	RETEVMO	SELPERCATINIB	LOXO ONCOLOGY INC	12/4/2019	5/8/2020	5.1	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER WITH ADVANCED OR METASTATIC RET FUSION-POSITIVE THYROID CANCER WHO REQUIRE SYSTEMIC THERAPY AND WHO ARE RADIOACTIVE IODINE-REFRACTORY (IF RADIOACTIVE IODINE-REFRACTORY (IF RADIOACTIVE IODINE).	Not Yet Converted	
NDA 213246	RETEVMO	SELPERCATINIB	LOXO ONCOLOGY INC	12/4/2019	5/8/2020	5.1	FOR THE TREATMENT OF ADULT PATIENTS WITH METASTATIC RET FUSION-POSITIVE NON-SMALL CELL LUNG CANCER (NSCLC)	Not Yet Converted	

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NDA 213246	RETEVMO	SELPERCATINIB	LOXO ONCOLOGY INC	12/4/2019	5/8/2020	5.1	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER WITH ADVANCED OR METASTATIC RET-MUTANT MEDULLARY THYROID CANCER (MTC) WHO REQUIRE SYSTEMIC THERAPY	Not Yet Converted	
NDA 213591	TABRECTA	CAPMATINIB	NOVARTIS PHARMACEUTICAL CORP	12/10/2019	5/6/2020	4.8	FOR THE TREATMENT OF ADULT PATIENTS WITH METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC) WHOSE TUMORS HAVE A MUTATION THAT LEADS TO MESENCHYMAL-EPITHELIAL TRANSITION (MET) EXON 14 SKIPPING AS DETECTED BY AN FDA-APPROVED TEST	Not Yet Converted	
BLA 125514 Supplement 59	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/18/2019	4/28/2020	12.4	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH	Not Yet Converted	
BLA 125514 Supplement 60	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/23/2019	4/28/2020	12.4	UNRESECTABLE OR METASTATIC MELANOMA PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH HEMATOLOGICAL MALIGNANCIES: HODGKIN LYMPHOMA	Not Yet Converted	
BLA 125514 Supplement 61	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/23/2019	4/28/2020	12.2	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH HEMATOLOGICAL MALIGNANCIES: PRIMARY MEDIASTINAL B-CELL LYMPHOMA	Not Yet Converted	
BLA 125514 Supplement 62	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/23/2019	4/28/2020	12.2	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATED WITH AN ANTI-ANGIOGENIC TYROSINE KINASE INHIBITOR (TKI)	Not Yet Converted	
BLA 125514 Supplement 63	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/23/2019	4/28/2020	12.2	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH MERKEL CELL CARCINOMA	Not Yet Converted	
BLA 125514 Supplement 64	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/23/2019	4/28/2020	12.2	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH ADVANCED OR METASTATIC GASTRIC OR GASTROESOPHAGEAL JUNCTION ADENOCARCINOMA	Not Yet Converted	
BLA 125514 Supplement 69	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	9/23/2019	4/28/2020	7.2	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH NON-SMALL CELL LUNG CANCER (NSCLC)	Not Yet Converted	
BLA 125514 Supplement 76	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/14/2020	4/28/2020	0.5	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH RENAL CELL CANCER (RCC)	Not Yet Converted	
BLA 125514 Supplement 77	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/14/2020	4/28/2020	0.5	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH ENDOMETRIAL CANCER	Not Yet Converted	
BLA 125514 Supplement 78	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/14/2020	4/28/2020	0.5	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH ADVANCED CERVICAL CANCER WITH DISEASE PROGRESSION DURING OR FOLLOWING CHEMOTHERAPY	Not Yet Converted	
BLA 125514 Supplement 79	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/14/2020	4/28/2020	0.5	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH BCG-UNRESPONSIVE, HIGH RISK, NON-MUSCLE INVASIVE BLADDER CANCER (NMIBC) WITH CARCINOMA IN-SITU (CIS) WITH OR WITHOUT PAPILLARY TUMORS WHO ARE INELIGIBLE FOR OR HAVE ELECTED NOT TO UNDERGO CYSTECTOMY	Not Yet Converted	
BLA 125514 Supplement 80	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/14/2020	4/28/2020	0.5	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH RECURRENT OR METASTATIC HEAD AND NECK SQUAMOUS CELL CARCINOMA (HNSCC) WITH DISEASE PROGRESSION ON OR AFTER PLATINUM-CONTAINING CHEMOTHERAPY	Not Yet Converted	
BLA 125514 Supplement 81	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/15/2020	4/28/2020	0.4	PROVIDES FOR AN ALTERNATE DOSE/SCHEDULE OF 400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH METASTATIC SCLC WITH DISEASE PROGRESSION ON OR AFTER PLATINUM-BASED CHEMOTHERAPY AND AT LEAST ONE OTHER LINE OF THERAPY	Not Yet Converted	

PROVIDES SOM AN ALTERNATE CODE/SIGNEDULE OF PROPERTY COLLARS AND ALTERNATE CODE SIGNEDULE OF PROPERTY COLLARS AND ALTERNATIVE COLLA						ppi ovais 200				
ROAD 241732 FEMALYITE PEMISHOLIZUMAB ROAD 241732 FEMALY PEMISHOLIZUMAB ROAD 241732 FEMALY PEMISHOLIZUMAB ROAD 241732 FEMALY PEMISHOLIZUMAB PEMISHOLIZU		KEYTRUDA	PEMBROLIZUMAB		4/15/2020	4/28/2020	0.4	RECURRENT LOCALLY ADVANCED OR METASTATIC ESOPHAGEAL CANCER WITH DISEASE PROGRESSION ON OR AFTER 2 OR MORE PRIOR LINES OF SYSTEMIC	Not Yet Converted	
BLA 125514 Superment 81 KEYTRUDA PEMBROLUZIMARA MERCK SHARP 8 DOWNER MERCK SHARP 8 MERCK S							***			
Bila 78115 TRODELYY SOCITIZIANAB SOCITIZIAN		KEYTRUDA	PEMBROLIZUMAB		4/20/2020	4/28/2020	0.3	400MG EVERY 6 WEEKS FOR ADULT PATIENTS WITH UNRESECTABLE OR METASTATIC MICROSATELLITE INSTABILITY-HIGH (MSI-H) OR MISMATCH REPAIR DEFICIENT (DMMR) SOLID TUMORS THAT HAVE PROGRESSED FOLLOWING PRIOR TX AND WHO HAVE	Not Yet Converted	
TREATED_UNRESSCTABLE LOCALLY ADVANCED OR METASTATIC CHALLANSOLAR CRISKING WITH A METASTATIC CHALLANSOLAR CRI	BLA 761115	TRODELVY		IMMUNOMEDICS INC	5/18/2018	4/22/2020	23.2	METASTATIC TRIPLE-NEGATIVE BREAST CANCER (MTNBC) WHO HAVE RECEIVED AT LEAST TWO PRIOR	Not Yet Converted	
NAZ 208574	NDA 213736	PEMAZYRE	PEMIGATINIB	INCYTE CORP	9/30/2019	4/17/2020	6.6	TREATED, UNRESECTABLE LOCALLY ADVANCED OR METASTATIC CHOLANGIOCARCINOMA WITH A FIBROBLAST GROWTH FACTOR RECEPTOR 2 (FGFR2) FUSION OR OTHER REARRANGEMENT AS DETECTED BY	Not Yet Converted	
BIL 125554 Supplement 78 OPDIVO NIVOLUMAB BRISTOL MYERS SQUIBB 9/10/2019 3/10/2020 6.0 TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATED WITH SORAFENIB NO OPDIVO NIVOLUMAB, FOR THE TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HOW THE PATIENT SWITH HOW THE PATIENT SWITH HOW THE PATIENT SWITH HOW BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HOW BEEN PREVIOUSLY TREATMENT OF PATIENTS WITH HOW THE PATIENT SWITH HOW BATTOR TO BUILT PATIENTS WITH HOW BATTOR TO BUILT PATIENTS WITH H			ROMIDEPSIN	PHARMACEUTICALS	8/18/2015	3/13/2020	54.9	LYMPHOMA (PTCL) IN ADULT PATIENTS WHO HAVE	Not Yet Converted	
BIA 125377 Supplement 108 YERVOY		OPDIVO	NIVOLUMAB		9/10/2019	3/10/2020	6.0	TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY	Not Yet Converted	
PATIENTS AGED 16 YEARS AND OLDER WITH METASTATIC OR IDCALLY ADVANCED EPITHELIOID Not Yet Converted		YERVOY	IPILIMUMAB		9/10/2019	3/10/2020	6.0	TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA (HCC) WHO HAVE BEEN PREVIOUSLY	Not Yet Converted	
BLA 761139 ENHERTU DAIICHI SANKYO INC 8/29/2019 12/20/2019 3.7 WIRESECTABLE OR METASTATIC HER2-POSITIVE BREAST CANCER WHO HAVE RECEIVED TWO OR MORE PRIOR ANTI-HER2-BASED REGIMENS IN THE NOT YET CONVERTED THE REATMENT OF ADULT PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CANCER (MUC) WHO HAVE PREVIOUSLY RECEIVED A PROGRAMMED DEATH-HIGAND 1 (PD-L1) INHIBITOR, AND A PLATINING CHEMOTHERAPY IN THE NEOADJUVANTI/ADJUVANT, LOCALLY ADVANCED OR PROGRAMMED DEATH-HIGAND 1 (PD-L1) INHIBITOR, AND A PLATINING CHEMOTHERAPY IN THE NEOADJUVANTI/ADJUVANT, LOCALLY ADVANCED OR METASTATIC SETTING BLA 761137 PADCEV VEDOTIAN-EJFV US INC 7/15/2019 12/18/2019 5.1 METASTATIC SETTING NOT YET CONVERTED THE NEOADJUVANTI/ADJUVANT, LOCALLY ADVANCED OR METASTATIC SETTING NOT YET CONVERTED THE NEOADJUVANTI/ADJUVANT, LOCALLY ADVANCED OR NOT YET CONVERTED THE NEOADJUVANTI/	NDA 211723	TAZVERIK	TAZEMETOSTAT	EPIZYME INC	5/23/2019	1/23/2020	8.1	PATIENTS AGED 16 YEARS AND OLDER WITH METASTATIC OR LOCALLY ADVANCED EPITHELIOID	Not Yet Converted	
LOCALLY ADVANCED OR METASTATIC UROTHELIAL CANCER (MUC) WHO HAVE PREVIOUSLY RECEIVED A PROGRAMMED DEATH RECEPTOR-1 (PD-1) OR PROGRAMMED DEATH-LIGAND 1 (PD-L1) INHIBITOR, AND A PLATINUM-CONTAINING CHEMOTHERAPY IN THE NEOADJUVANT/ADJUVANT, LOCALLY ADVANCED OR Not Yet Converted SAREPTA NDA 211970 VYONDYS 53 GOLODIRSEN THERAPEUTICS INC 12/19/2018 12/12/2019 11.8 AMENABLE TO EXON 33 SKIPPING NDA 213137 OXBRYTA VOXELOTOR THERAPEUTICS INC 6/26/2019 11/25/2019 5.0 AND OLDER FOR THE TREATMENT OF FADULT PATIENTS WITH AND LOCALLY ADVANCED OR REVIVED A PROGRAMMED DEATH-LIGAND 1 (PD-L1) INHIBITOR, AND A PLATINUM-CONTAINING CHEMOTHERAPY IN THE NEOADJUVANT/ADJUVANT, LOCALLY ADVANCED OR Not Yet Converted FOR THE TREATMENT OF DUCHENNE MUSCULAR CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 33 SKIPPING Not Yet Converted FOR THE TREATMENT OF SICKLE CELL DISEASE IN ADULTS AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER FOR THE TREATMENT OF ADULT PATIENTS WITH MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED	BLA 761139	ENHERTU	TRASTUZUMAB DERUXTECAN-	DAIICHI SANKYO INC	8/29/2019	12/20/2019	3.7	UNRESECTABLE OR METASTATIC HER2-POSITIVE BREAST CANCER WHO HAVE RECEIVED TWO OR MORE PRIOR ANTI-HER2-BASED REGIMENS IN THE	Not Yet Converted	
NDA 211970 VYONDYS 53 GOLODIRSEN SAREPTA NDA 211970 VYONDYS 53 GOLODIRSEN THERAPEUTICS INC 12/19/2018 12/12/2019 11.8 AMENABLE TO EXON 53 SKIPPING NDA 213137 OXBRYTA VOXELOTOR THERAPEUTICS INC 6/26/2019 11/25/2019 5.0 AND OLDER FOR THE TREATMENT OF DUCHENNE MUSCULAR DYSTROPHY (MDM) IN PATIENTS WHO HAVE A CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 53 SKIPPING FOR THE TREATMENT OF DUCHENNE MUSCULAR DYSTROPHY (MDM) IN PATIENTS WHO HAVE A CONFIRMED MUSCULAR DYSTROPHY (MDD) IN PATIENTS WHO HAVE A CONFIRMED MUSCULAR DYSTR	RI & 761127	PADCEV			7/15/2010	12/18/2019	51	LOCALLY ADVANCED OR METASTATIC UROTHELIAL CANCER (MUC) WHO HAVE PREVIOUSLY RECEIVED A PROGRAMMED DEATH RECEPTOR-1 (PD-1) OR PROGRAMMED DEATH-LIGAND 1 (PD-1) INHIBITOR, AND A PLATINUM-CONTAINING CHEMOTHERAPY IN THE NEOADJUVANT/ADJUVANT, LOCALLY ADVANCED OR	Not Vet Converted	
NDA 211970 VYONDYS 53 GOLODIRSEN THERAPEUTICS INC 12/19/2018 12/12/2019 11.8 DYSTROPHY (DMD) IN PATIENTS WHO HAVE A CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 53 SKIPPING NOT YET CONVERTED WITH THE TRAPEUTICS INC 12/19/2018 12/12/2019 11.8 AMENABLE TO EXON 53 SKIPPING NOT YET CONVERTED WITH THE TRAPEUTICS INC 6/26/2019 11/25/2019 5.0 AND OLDER NOT YET CONVERTED NOT YET CONVERTED WITH MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED	BLA /6113/	PADCEV	VEDUTIAN-EJFV	US INC	7/15/2019	12/18/2019	5.1		Not yet Converted	
NDA 213137 OXBRYTA VOXELOTOR THERAPEUTICS INC 6/26/2019 11/25/2019 5.0 AND OLDER Not Yet Converted FOR THE TREATMENT OF ADULT PATIENTS WITH MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED	NDA 211970	VYONDYS 53	GOLODIRSEN	THERAPEUTICS INC	12/19/2018	12/12/2019	11.8	DYSTROPHY (DMD) IN PATIENTS WHO HAVE A CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 53 SKIPPING FOR THE TREATMENT OF SICKLE CELL DISEASE IN	Not Yet Converted	
MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED	NDA 213137	OXBRYTA	VOXELOTOR		6/26/2019	11/25/2019	5.0	AND OLDER	Not Yet Converted	
	NDA 213217	BRUKINSA	ZANUBRUTINIB	BEIGENE USA INC	6/27/2019	11/14/2019	4.6	MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED	Not Yet Converted	

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BLA 125514 Supplement 65	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	6/17/2019	9/17/2019	3.0	IN COMBINATION WITH LENVATINIB, INDICATED FOR THE TREATMENT OF PATIENTS WITH ADVANCED ENDOMETRIAL CARCINOMA THAT IS NOT MICROSATELLITE INSTABILITY-HIGH (MSI-H) OR MISMATCH REPAIR DEFICIENT (DMMR), WHO HAVE DISEASE PROGRESSION FOLLOWING PRIOR SYSTEMIC THERAPY AND ARE NOT CANDIDATES FOR CURATIVE SURGERY OR RADIATION	Not Yet Converted	
NDA 206947							IN COMBINATION WITH PEMBROLIZUMAB, INDICATED FOR THE TREATMENT OF PATIENTS WITH ADVANCED ENDOMETRIAL CARCINOMA THAT IS NOT MICROSATELLITE INSTABILITY-HIGH (MSI-H) OR MISMATCH REPAIR DEFICIENT (DMMR), WHO HAVE DISEASE PROGRESSION FOLLOWING PRIOR SYSTEMIC THERAPY AND ARE NOT CANDIDATES FOR CURATIVE		
Supplement 11 NDA 212726 Original 1	LENVIMA ROZLYTREK	LENVATINIB	EISAI INC	6/17/2019	9/17/2019 8/15/2019	7.9	SURGERY OR RADIATION FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER WITH SOLID TUMORS THAT HAVE A NEUROTROPHIC TYROSINE RECEPTOR KINASE (NTRK) GENE FUSION WITHOUT A KNOWN ACQUIRED RESISTANCE MUTATION, ARE METASTATIC OR WHERE SURGICAL RESECTION IS LIKELY TO RESULT IN SEVER MORBIDITY, AND HAVE PROGRESSED FOLLOWING TREATMENT OR HAVE SATISFACTORY ALTERNATIVE THERAPY	Not Yet Converted Not Yet Converted	
NDA 204384 Supplement 10	SIRTURO	BEDAQUILINE	JANSSEN RESEARCH AND DEVELOPMENT LLC	10/11/2018	8/9/2019	9.9	AS PART OF COMBINATION THERAPY FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS (12 TO LESS THAN 18 YEARS OF AGE AND WEIGHING AT LEAST 30KG) WITH PULMONARY MULTI-DRUG RESISTANT TUBERCULOSIS (MDR-TB)	Not Yet Converted	
NDA 212306	XPOVIO	SELINEXOR	KARYOPHARM THERAPEUTICS INC	8/6/2018	7/3/2019	10.9	IN COMBINATION WITH DEXAMETHASONE FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY MULTIPLE MYELOMA (RRMM) WHO HAVE RECEIVED AT LEAST FOUR PRIOR THERAPIES AND WHOSE DISEASE IS REFRACTORY TO AT LEAST TWO PROTEASOME INHIBITORS, AT LEAST TWO IMMUNOMODULATORY AGENTS, AND AN ANTI-CD38 MONOCLONAL ANTIBODY	Converted	12/18/2020
BLA 125514 Supplement 53	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	12/17/2018	6/17/2019	6.0	FOR THE TREATMENT OF PATIENTS WITH METASTATIC SMALL CELL LUNG CANCER WITH DISEASE PROGRESSION ON OR AFTER PLATINUM-BASED CHEMOTHERAPY AND AT LEAST ONE OTHER PRIOR LINE OF THERAPY	Not Yet Converted	12102020
BLA 761121	POLIVY	POLATUZUMAB VEDOTIN-PIIQ	GENENTECH	12/19/2018	6/10/2019	5.7	IN COMBINATION WITH BENDAMUSTINE AND RITUXIMAB PRODUCT, FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY DIFFUSE LARGE B-CELL LYMPHOMA, NOT OTHERWISE SPECIFIED, AFTER AT LEAST TWO PRIOR THERAPIES	Not Yet Converted	
NDA 212018	BALVERSA	ERDAFITINIB	JANSSEN BIOTECH	9/18/2018	4/12/2019	6.8	FOR THE TREATMENT OF ADULT PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (MUC), THAT HAS: SUSCEPTIBLE FGFR3 OR FGFR2 GENETIC ALTERATIONS AND PROGRESSED DURING OR FOLLOWING AT LEAST ONE LINE OF PRIOR PLATINUM CONTAINING CHEMOTHERAPY, INCLUDING WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT PLATINUM CONTAINING CHEMOTHERAPY	Not Yet Converted	
BLA 761034 Supplement 18	TECENTRIQ	ATEZOLIZUMAB	GENENTECH	9/12/2018	3/8/2019	5.8	IN COMBINATION WITH PACLITAXEL PROTEIN-BOUND FOR THE TREATMENT OF ADULT PATIENTS WITH UNRESECTABLE LOCALLY ADVANCED OR METASTATIC TRIPLE-NEGATIVE BREAST CANCER (TNBC) WHOSE TUMORS EXPRESS PD-L1 (PD-L1 STAINED TUMOR-INFILTRATING IMMUNE CELLS [IC] OF ANY INTENSITY COVERING 2 1% OF THE TUMOR AREA), AS DETERMINED BY AN FDA-APPROVED TEST	Not Yet Converted	
BLA 125514 Supplement 45	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	6/29/2018	12/19/2018	5.7	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS WITH RECURRENT LOCALLY ADVANCED OR METASTATIC MERKEL CELL CARCINOMA	Not Yet Converted	

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NDA 210861	VITRAKVI (CAP SULES)	LAROTRECTINIB	LOXO ONCOLOGY	3/26/2018	11/26/2018	8.1	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS WITH SOLID TUMORS THAT HAVE A NEUROTROPHIC RECEPTOR TYROSINE KINASE (NTRK) GENE FUSION WITHOUT A KNOWN ACQUIRED RESISTANCE MUTATION; ARE METASTATIC OR WHERE SURGICAL RESECTION IS LIKELY TO RESULT IN SEVERE MORBIDITY; AND HAVE NO SATISFACTORY ALTERNATIVE TREATMENTS OR THAT HAVE PROGRESSED FOLLOWING TREATMENT FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS WITH SOLID TUMORS THAT HAVE A NEUROTROPHIC RECEPTOR TYROSINE KINASE (NTRK) GENE FUSION WITHOUT A KNOWN ACQUIRED RESISTANCE MUTATION; ARE METASTATIC OR WHERE SURGICAL RESECTION IS LIKELY TO RESULT IN SEVERE MORBIDITY; AND HAVE NO SATISFACTORY ALTERNATIVE TREATMENTS OR THAT HAVE	Not Yet Converted	
NDA 211710 NDA 208573 Supplement 9	L SOLUTION) VENCLEXTA	VENETOCLAX	LOXO ONCOLOGY ABBVIE	3/26/2018 6/25/2018	11/26/2018	8.1	PROGRESSED FOLLOWING TREATMENT VENCLEXTA-IN COMBINATION WITH AZACITIDINE OR DECITABINE OR LOW-DOSE CYTARABINE FOR THE TREATMENT OF NEWLY-DIAGNOSED ACUTE MYELOID LEUKEMIA (AML) IN ADULTS WHO ARE AGE 75 YEARS OR OLDER, OR WHO HAVE COMORBIDITIES THAT PRECLUDE USE OF INTENSIVE INDUCTION CHEMOTHERAPY	Not Yet Converted Converted	5/22/2020
BLA 125514 Supplement 42	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	5/9/2018	11/9/2018	6.0	FOR THE TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA WHO HAVE BEEN PREVIOUSLY TREATED WITH SORAFENIB	Not Yet Converted	
NDA 210868	LORBRENA	LORLATINIB	PFIZER	12/5/2017	11/2/2018	10.9	FOR THE TREATMENT OF PATIENTS WITH ANAPLASTIC LYMPHOMA KINASE (ALK)-POSITIVE METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC) WHOSE DISEASE HAS PROGRESSED ON: • CRIZOTINIB AND AT LEAST ONE OTHER ALK INHIBITOR FOR METASTATIC DISEASE; OR • ALECTINIB AS THE FIRST ALK INHIBITOR THERAPY FOR METASTATIC DISEASE; OR • CERITINIB AS THE FIRST ALK INHIBITOR THERAPY FOR METASTATIC DISEASE; OR	Not Yet Converted	
NDA 207356	ARIKAYCE	AMIKACIN LIPOSOME INHALATION SUSPENSION	INSMED	3/28/2018	9/28/2018	6.0	ARIKAYCE IS AN AMINOGLYCOSIDE ANTIBACTERIAL INDICATED IN ADULTS WHO HAVE LIMITED OR NO ALTERNATIVE TREATMENT OPTIONS, FOR THE TREATMENT OF MYCOBACTERIUM AVIUM COMPLEX (MAC), LUNG DISEASE AS PART OF A COMBINATION ANTIBACTERIAL DRUG REGIMEN IN PATIENTS WHO DO NOT ACHIEVE NEGATIVE SPUTUM CULTURES AFTER A MINIMUM OF 6 CONSECUTIVE MONTHS OF A MULTIDRUG BACKGROUND REGIMEN THERAPY. AS ONLY LIMITED CLINICAL SAFETY AND EFFECTIVENESS DATA FOR ARIKAYCE FOR USE IN ADULTS WHO HAVE LIMITED OR NO ALTERNATIVE TREATMENT OPTIONS. THIS DRUG IS INDICATED FOR USE IN A LIMITED AND SPECIFIC POPULATION OF PATIENTS. THIS INDICATION IS APPROVED UNDER ACCELERATED APPROVAL BASED ON ACHIEVING SPUTUM CULTURE CONVERSION (DEFINED AS 3 CONSECUTIVE NEGATIVE MONTHLY SPUTUM CULTURES) BY MONTH 6. CLINICAL BENEFIT HAS NOT YET BEEN ESTABLISHED	Not Yet Converted	
NDA 211155	COPIKTRA	DUVELISIB	VERASTEM	2/5/2018	9/24/2018	7.6	FOR THE TREATMENT OF ADULT PATIENTS WITH RELAPSED OR REFRACTORY FOLLICULAR LYMPHOMA (FL) AFTER AT LEAST TWO PRIOR SYSTEMIC THERAPIES	Not Yet Converted	
BLA 125554 Supplement 67	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	2/16/2018	8/16/2018	6.0	FOR THE TREATMENT OF PATIENTS WITH METASTATIC SMALL CELL LUNG CANCER (SCLC) WITH PROGRESSION AFTER PLATINUM-BASED CHEMOTHERAPY AND AT LEAST ONE OTHER LINE OF THERAPY	Withdrawn	12/29/2020

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NDA 208623	GALAFOLD	MIGALASTAT	AMICUS THERAPEUTICS	12/13/2017	8/10/2018	7.9	FOR THE TREATMENT OF ADULTS WITH A CONFIRMED DIAGNOSIS OF FABRY DISEASE AND AN AMENABLE GALACTOSIDASE ALPHA GENE (GLA) VARIANT BASED ON IN VITRO ASSAY DATA	Not Yet Converted	
BLA 125554 Supplement 63	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	1/10/2018	7/10/2018	6.0	IN COMBINATION WITH IPILIMUMAB FOR THE TREATMENT OF ADULTS AND PEDIATRIC PATIENTS 12 YEARS AND OLDER WITH MICROSATELLITE INSTABLITY- HIGH (MSI H) OR DNA MISMATCH REPAIR DEFICIENT (DMMR), METASTATIC COLORECTAL CANCER THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMIDINE, OXALIPLATIN, AND IRINOTECAN	Not Yet Converted	
BLA 125377 Supplement 96	YERVOY	IPILIMUMAB	BRISTOL MYERS SQUIBB	1/10/2018	7/10/2018	6.0	IN COMBINATION WITH NIVOLUMAB FOR THE TREATMENT OF ADULTS AND PEDIATRIC PATIENTS 12 YEARS AND OLDER WITH MICROSATELLITE INSTABILITY- HIGH (MSI H) OR DNA MISMATCH REPAIR DEFICIENT (DMMR), METASTATIC COLORECTAL CANCER THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMIDINE, OXALIPLATIN, AND IRINOTECAN	Not Yet Converted	
Supplement 90	TERVOT	IFILINOWAB	SQUIBB	1/10/2016	7/10/2018	0.0	FOR THE TREAMENT OF ADULT AND PEDIATRIC	Not ret Convented	
BLA 125514 Supplement 30	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	10/3/2017	6/13/2018	8.3	PATIENTS WITH REFRACTORY PRIMARY MEDIASTINAL LARGE B-CELL LYMPHOMA (PMBCL), OR WHO HAVE RELAPSED AFTER 2 OR MORE PRIOR LINES OF THERAPY	Converted	10/14/2020
BLA 125514 Supplement 34	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	12/28/2017	6/12/2018	5.5	FOR THE TREATMENT OF PATIENTS WITH RECURRENT OR METASTATIC CERVICAL CANCER WITH DISEASE PROGRESSION ON OR AFTER CHEMOTHERPAY WHOSES TUMORS EXPRESS PD-L1 (CPS > 1) AS DETERMINED BY AN FDA-APPROVED TEST	Not Yet Converted	
NDA 21462 Supplement 51	ALIMTA	PEMETREXED DISODIUM	ELI LILLY AND COMPANY	8/3/2017	6/4/2018	10.0	IN COMBINATION WITH PEMBROLIZUMAB AND CARBOPLATIN, FOR THE FIRST-LINE TREATMENT OF METASTATIC, NON-SQUAMOUS, NON-SMALL CELL LUNG CANCER	Converted	1/30/2019
BLA 125557 Supplement 13	BLINCYTO	BLINATUMOMAB	AMGEN	9/29/2017	3/29/2018	6.0	FOR THE TREAMENT OF B-CELL PRECURSOR ACUTE LYMPHOBLASTIC LEUKEMIA (ALL) IN FIRST OR SECOND COMPLETED REMISSION WITH MINIMAL RESIDUAL DISEASE (MRD) GREATER OR EQUAL TO 0.1% IN ADULTS AND CHILDREN	Not Yet Converted	
NDA 210563 Original 2	IMBRUVICA	IBRUTINIB	PHARMACYCLICS	8/31/2017	2/16/2018	5.6	FOR THE TREATMENT OF ADULT PATIENTS WITH MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY	Not Yet Converted	
NDA 210563 Original 2	IMBRUVICA	IBRUTINIB	PHARMACYCLICS	8/31/2017	2/16/2018	5.6	FOR THE TREATMENT OF ADULT PATIENTS WITH MARGINAL ZONE LYMPHOMA (MZL) WHO REQUIRE SYSTEMIC THERAPY AND HAVE RECEIVED AT LEAST ONE PRIOR ANTI- CD20 BASED THERAPY	Not Yet Converted	
NDA 203341 Supplement 9	BOSULIF	BOSUTINIB	PF PRISM CV	6/26/2017	12/19/2017	5.8	FOR THE TREATMENT OF ADULT PATIENTS WITH NEWLY DIAGNOSED CHRONIC PHASE (CP) PHILADELPHIA CHROMOSOME POSITIVE CHRONIC MYELOGENOUS LEUKEMIA (PH+ CML)	Not Yet Converted	
NDA 210259	CALQUENCE	ACALABRUTINIB	ASTRAZENECA PHARMACEUTICALS	6/13/2017	10/31/2017	4.6	FOR THE TREATMENT OF ADULT PATIENTS WITH MANTLE CELL LYMPHOMA (MCL) WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY	Not Yet Converted	
BLA 125554 Supplement 41	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	3/24/2017	9/22/2017	6.0	FOR THE TREATMENT OF PATIENTS WITH HEPATOCELLULAR CARCINOMA WHO HAVE BEEN PREVIOUSLY TREATED WITH SORAFENIB	Not Yet Converted	

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ALICOPA		KEYTRUDA	PEMBROLIZUMAB		3/22/2017	9/22/2017	6.0	LOCALLY ADVANCED OR METASTATIC GASTRIC OR GASTROESOPHAGEAL JUNCTION ADENOCARCINOMA WHOSE TUMORS EXPRESS PD-L1 (COMBINED POSITIVE SCORE (CPS) 21] AS DETERMINED BY AN FDA-APPROVED TEST, WITH DISEASE PROGRESSION ON OR AFTER TWO OR MORE PRIOR LINES OF THERAPY INCLUDING FLUOROPYRIMIDINE- AND PLATINUM-CONTAINING CHEMOTHERAPY AND IF APPROPRIATE,	Not Yet Converted	
CHEMO RESEARCH AL CO EXCELTS 1292016 8292017 8.0 FOR THE TREATMENT OF CHAGAS DISEASE (AMERICA) Not Yet Convented						0/11/0017		RELAPSED FOLLICULAR LYMPHOMA (FL) WHO HAVE		
CHEMO RESEARCH 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2016 1229/2017 1229/2016 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017 1229/2017 1229/2016 1229/2017 1229/2016 1229/2017	NDA 209936	ALIQOPA	COPANLISIB	PHARMACEUTICALS	3/16/2017	9/14/2017	6.0		Not Yet Converted	
BLA 125514 REYTRUDA PEMBROLIZUMAB BRISTOL MYES SUBJOINT	NDA 209570	N/A ¹	BENZNIDAZOLE	AL C/O EXCELTIS	12/29/2016	8/29/2017	8.0	TRYPANOSOMIASIS), CAUSED BY TRYPANOSOMA CRUZI,	Not Yet Converted	
PATIENTS WITH UNRESECTABLE OR METASTATIC, MICROSATELLITE MISSHIP OR METASTATIC, MICROSATELLITE MISSHIP OR MISSHATOR REPAIR DEFICIENT SOLID TUMORS THAT HAVE PROGRESSED FOLLOWING PROOF THEATMENT AND WHO HAVE NO SATISFACTORY ALTERNATIVE THOU THEATMENT OF CHRONIC IRON OF MISSHATORY REPAIR DEFICIENT COLORECTAL CANCER THAT HAS PROGRESSED FOLLOWING PROOF TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED A PLANTAGE OF THE TREATMENT OF CHRONIC IRON OVERLOAD UNDER THAT HAS PROGRESSED FOLLOWING PROOF THE AND THAT HAS PROGRESSED FOLLOWING PROOF THE AND THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED AND THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED AND THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED AND THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED AND THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMDINE, OXALPLATIN, AND RINDTECAN NOT YEL CONVERTED AND THAT HAS PROBLED AND THAT H		OPDIVO	NIVOLUMAB		2/2/2017	7/31/2017	5.9	PATIENTS 12 YEARS AND OLDER WITH MICROSATELLITE INSTABILITY-HIGH (MSI-H) OR MISMATCH REPAIR DEFICIENT (DMMR) METASTATIC COLORECTAL CANCER (CRC) THAT HAS PROGRESSED FOLLOWING TREATMENT WITH A FLUOROPYRIMIDINE, OXALIPLATIN,	Not Yet Converted	
NOVARTIS		KEYTRUDA	PEMBROLIZIMAB		9/8/2016	5/23/2017	54	PATIENTS WITH UNRESECTABLE OR METASTATIC, MICROSATELLITE INSTABILITY-HIGH (MSI-H) OR MISMATCH REPAIR DEFICIENT SOLID TUMORS THAT HAVE PROGRESSED FOLLOWING PRIOR TREATMENT AND WHO HAVE NO SATISFACTORY ALTERNATIVE TREATMENT OPTIONS, OR METASTATIC MICROSATELLITE INSTABILITY-HIGH (MSH-H) OR MISMATCH REPAIR DEFICIENT COLORECTAL CANCER THAT HAS PROGRESSED FOLLOWING TREATMENT WITH	Not Yet Converted	
PATIENTS 10 YEARS OF AGE AND DIDER WITH NON- TRANSFUSION-DEPENDENT THALASSEMIA (NTDT) SYNDROMES AND WITH A LIVER IRON CONCENTRATION (LIC) OF AT LEAST 5 MILLIGARMS OF IRON PER GRAM OF LIVER DRY WEIGHT (MG FE/G DW) AND A SERUM FERRITIN GREATER THAN 300 MCG/L BLA 125514 Supplement 17 BLA 125514 Supplement 16 KEYTRUDA MERCK SHARP & DOHME MERCK SHARD MONTHS DEAST SWITH LOCALLY MONTHS DEAST SWITH LOCALLY		JADENU		NOVARTIS				DUE TO BLOOD TRANSFUSIONS (TRANSFUSIONAL HEMOSIDEROSIS) IN PATIENTS 2 YEARS OF AGE AND		
BLA 125514 Supplement 17 KEYTRUDA PEMBROLIZUMAB MERCK SHARP & DOHME 12/14/2016 5/18/2017 5.1 FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO ARE NOT ELIGIBLE FOR CISPLATIN-CONTAINING CHEMOTHERAPY IN COMBINATION WITH PEMETREXED AND CARBOPLATIN, FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH METASTATIC NON-SQUAMOUS, NON-SMALL CELL LUING CANCER FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (UC) WHO: HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY		JADENU		NOVARTIS				FOR THE TREATMENT OF CHRONIC IRON OVERLOAD IN PATIENTS 10 YEARS OF AGE AND OLDER WITH NON-TRANSFUSION-DEPENDENT THALASSEMIA (NTDT) SYNDROMES AND WITH A LIVER IRON CONCENTRATION (LIC) OF AT LEAST 5 MILLIGRAMS OF IRON PER GRAM OF LIVER DRY WEIGHT (MG FE/G DW) AND A SERUM		
BLA 125514 Supplement 17 BLA 125514 Supplement 17 BLA 125514 Supplement 16 KEYTRUDA PEMBROLIZUMAB MERCK SHARP & DOHME DOHME DOHME DOHME 12/14/2016 5/18/2017 5.1 ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO ARE NOT ELIGIBLE FOR CISPLATIN-CONTAINING CHEMOTHERAPY IN COMBINATION WITH PEMETREXED AND CARBOPLATIN, FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH METASTATIC NON-SQUAMOUS, NON-SMALL CELL LUNG CANCER FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (UC) WHO: HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTH OF ADJUVANT OR ADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY	NDA 207968	SPRINKLE	DEFERASIROX	PHARMACEUTICALS	7/21/2016	5/18/2017	9.9		Converted	7/23/2020
BLA 125514 Supplement 16 KEYTRUDA PEMBROLIZUMAB MERCK SHARP & DOHME 11/10/2016 5/10/2017 6.0 SMALL CELL LUNG CANCER FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH METASTATIC UNCN-SQUAMOUS, NON- SMALL CELL LUNG CANCER FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (UC) WHO: HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY WITH PLATINUM-		KEYTRUDA	PEMBROLIZUMAB		12/14/2016	5/18/2017	5.1	ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO ARE NOT ELIGIBLE FOR CISPLATIN-CONTAINING CHEMOTHERAPY	Not Yet Converted	
FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (UC) WHO: HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY WITH PLATINUM-CONTAINING CHEMOTHERAPY		KEYTRUDA	PEMBROLIZUMAR		11/10/2016	5/10/2017	6.0	CARBOPLATIN, FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH METASTATIC NON-SQUAMOUS, NON-	Converted	8/20/2018
								FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA (UC) WHO: HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR WHO HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT TREATMENT		

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BLA 761069	IMFINZI	DURVALUMAB	ASTRAZENECA PHARMACEUTICALS	10/13/2016	5/1/2017	6.6	FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY	Not Yet Converted	
NDA 208772	ALUNBRIG	BRIGATINIB	ARIAD PHARMACEUTICALS	8/29/2016	4/28/2017	8.0	FOR TREATMENT OF PATIENTS WITH ANAPLASTIC LYMPHOMA KINASE (ALK)-POSITIVE METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC) WHO HAVE PROGRESSED ON OR ARE INTOLERANT TO CRIZOTINIB	Converted	5/22/2020
BLA 125554 Supplement 31	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	12/9/2016	4/25/2017	4.5	FOR THE TREATMENT OF ADULT PATIENTS WITH CLASSICAL HODGKIN LYMPHOMA THAT HAS RELAPSED OR PROGRESSED AFTER: * AUTOLOGOUS HEMATOPOIETIC STEM CELL TRANSPLANTATION (HSCT) AND BRENTUXIMAB VEDOTIN, OR *3 OR MORE LINES OF SYSTEMIC THERAPY THAT INCLUDES AUTOLOGOUS HSCT.	Not Yet Converted	
BLA 761034 Supplement 1	TECENTRIQ	ATEZOLIZUMAB	GENENTECH	10/31/2016	4/17/2017	5.5	PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO ARE NOT ELIGIBLE FOR CISPLATIN-CONTAINING CHEMOTHERAPY, OR HAVE DISEASE PROGRESSION DURING OR FOLLOWING ANY PLATINUM-CONTAINING CHEMOTHERAPY, OR WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT CHEMOTHERAPY	Not Yet Converted	
BLA 761049	BAVENCIO	AVELUMAB	EMD SERONO	9/23/2016	3/23/2017	6.0	FOR THE TREATMENT OF ADULTS AND PEDIATRIC PATIENTS 12 YEARS AND OLDER WITH METASTATIC MERKEL CELL CARCINOMA	Not Yet Converted	
BLA 125514 Supplement 15	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	9/15/2016	3/14/2017	5.9	FOR THE TREATMENT OF ADULT AND PEDIATRIC PATIENTS WITH REFRACTORY CLASSICAL HODGKIN LYMPHOMA, OR WHO HAVE RELAPSED AFTER 3 OR MORE PRIOR LINES OF THERAPY	Converted	10/14/2020
BLA 125554 Supplement 24	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	9/2/2016	2/2/2017	5.0	FOR THE TREATMENT OF LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO: • HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY • HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY.	Not Yet Converted	
NDA 205552 Supplement 16	IMBRUVICA	IBRUTINIB	PHARMACYCLICS	9/23/2016	1/18/2017	3.8	FOR THE TREATMENT OF PATIENTS WITH MARGINAL ZONE LYMPHOMA (MZL.) WHO REQUIRE SYSTEMIC THERAPY AND HAVE RECEIVED AT LEAST ONE PRIOR ANTI-CD20-BASED THERAPY	Not Yet Converted	
NDA 000445	DUDDAGA	BUGABARIA		0/00/0040	40/40/0040	5.0	FOR THE TREATMENT OF PATIENTS WITH DELETERIOUS BRCA MUTATION (GERMLINE AND/OR SOMATIC) ASSOCIATED ADVANCED OVARIAN CANCER WHO HAVE BEEN TREATED WITH TWO OR MORE CHEMOTHERAPIES	2	4/0/0040
NDA 209115	RUBRACA	RUCAPARIB	CLOVIS ONCOLOGY ELI LILLY AND	6/23/2016	12/19/2016	5.9	IN COMBINATION WITH DOXORUBICIN FOR THE TREATMENT OF ADULT PATIENTS WITH SOFT TISSUE SARCOMA (STS) WITH A HISTOLOGIC SUBTYPE FOR WHICH AN ANTHRACYCLINE-CONTAINING REGIMEN IS APPROPRIATE AND WHICH IS NOT AMENABLE TO CURATIVE TREATMENT WITH RADIOTHERAPY OR	Converted	4/6/2018
BLA 761038	LARTRUVO	OLARATUMAB	COMPANY	2/24/2016	10/19/2016	7.8	SURGERY	Not Yet Converted	
NDA 206488	EXONDYS 51	ETEPLIRSEN	SAREPTA THERAPEUTICS	6/26/2015	9/19/2016	14.8	FOR THE TREATMENT OF DUCHENNE MUSCULAR DYSTROPHY (DMD) IN PATIENTS WHO HAVE A CONFIRMED MUTATION OF THE DMD GENE THAT IS AMENABLE TO EXON 51 SKIPPING	Not Yet Converted	
BLA 125514	WENTEN !	DEMODOLIZIE:::	MERCK SHARP &	0/0/0046	0/5/0040	50	FOR THE TREATMENT OF PATIENTS WITH RECURRENT OR METASTATIC SQUAMOUS CELL CARCINOMA OF THE HEAD AND NECK WITH DISEASE PROGRESSION ON OR AFTER PLATINUM-CONTAINING CHEMOTHERAPY	Quantum .	
Supplement 9	KEYTRUDA	PEMBROLIZUMAB	DOHME	2/9/2016	8/5/2016	5.9		Converted	6/10/2019

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		OBETICHOLIC	INTERCEPT				FOR THE TREATMENT OF PRIMARY BILIARY CHOLANGITIS (PBC) IN COMBINATION WITHURSODEOXYCHOLIC ACID (UDCA) IN ADULTS WITH AN INADEQUATE RESPONSE TO UDCA, OR AS MONOTHERAPY IN ADULTS UNABLE TO TOLERATE UDCA		
NDA 207999	OCALIVA	ACID	PHARMACEUTICALS	6/29/2015	5/27/2016	10.9		Not Yet Converted	
							FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC UROTHELIAL CARCINOMA WHO HAVE DISEASE PROGRESSION DURING OR FOLLOWING PLATINUM-CONTAINING CHEMOTHERAPY OR HAVE DISEASE PROGRESSION WITHIN 12 MONTHS OF NEOADJUVANT OR ADJUVANT TREATMENT WITH PLATINUM-CONTAINING CHEMOTHERAPY		
BLA 761034	TECENTRIQ	ATEZOLIZUMAB	GENENTECH	1/12/2016	5/18/2016	4.2		Not Yet Converted	
BLA 125554 Supplement 19	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	3/1/2016	5/17/2016	2.5	FOR THE TREATMENT OF CLASSICAL HODGKIN LYMPHOMA THAT HAS RELAPSED OR PROGRESSED AFTER AUTOLOGOUS HEMATOPOLETIC STEM CELL TRANSPLANTATION (HSCT), AND POST-	Not Yet Converted	
Supplement 19	OFDIVO	INIVOLUMAB	SQUIBB	3/1/2010	3/17/2010	2.5	TRANSPLANTATION BRENTUXIMAB VEDOTIN. FOR THE TREATMENT OF PATIENTS WITH CHRONIC	Not ret Convented	
NDA 208573	VENCLEXTA	VENETOCLAX	ABBVIE	10/29/2015	4/11/2016	5.4	LYMPHOCYTIC LEUKEMIA (CLL) WITH 17P DELETION, AS DETECTED BY AN FDA APPROVED TEST, WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY	Converted	6/8/2018
						-	FOR THE TREATMENT OF PEDIATRIC AND ADULT		
NDA 204630	PROVAYBLUE	METHYLENE BLUE	PROVEPHARM SAS	10/9/2015	4/8/2016	6.0	PATIENTS WITH ACQUIRED METHEMOGLOBINEMIA 1) IN COMBINATION WITH IPILIMUMAB FOR THE TREATMENT OF PATEINTS WITH UNRESECTABLE OR METASTATIC MELANOMA TO REMOVE THE RESTRICTION FOR THE TREATMENT OF ONLY PATIENTS WIHT BRAF WILD-TYPE MELANOMA, 2) AS A SINGLE A GENT FOR THE TREATMENT OF PATIENTS WITH BRAF V600 MUTATION POSITIVE, UNRESECTABLE OR METASTATIC MELANOMA TO REMOVE THE RESTRICTION THAT SUCH PATIENTS SHOULD HAVE DISEASE PROGRESSION	Not Yet Converted	
BLA 125554 Supplement 7	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	7/23/2015	1/23/2016	6.0	FOLLOWING IPILIMUMAB AND A BRAF INHIBITOR	Converted	3/7/2019
NDA 208434	ALECENSA	ALECTINIB	HOFFMAN LA ROCHE				FOR THE TREATMENT OF PATIENTS WITH ANAPLASTIC LYMPHOMA KINASE (ALK)-POSITIVE METASTATIC NON-SMALL CELL LUNG CANCER (NSCLO), WHO HAVE PROGRESSED ON OR ARE INTOLERANT TO CRIZOTINIB		11/6/2017
NDA 206434	ALECENSA	ALECTINIB	HOFFMAN LA ROCHE	7/6/2015	12/11/2015	5.2	FOR TREATMENT OF PATIENTS WITH MULTIPLE	Converted	11/6/2017
BLA 761036	DARZALEX	DARATUMUMAB	JANSSEN BIOTECH	7/9/2015	11/16/2015	4.3	MYELOMA WHO HAVE RECEIVED AT LEAST 3 PRIOR LINES OF THERAPY INCLUDING A PROTEASOME INHIBITOR AND AN IMMUNOMODULATORY AGENT OR ARE DOUBLE REFRACTORY TO A PROTEASOME INHIBITOR AND AN IMMUNOMODULATORY AGENT	Converted	11/21/2016
NDA 208065	TAGRISSO	OSIMERTINIB	ASTRAZENECA PHARMACEUTICALS	6/5/2015	11/13/2015	5.3	FOR THE TREATMENT OF PATIENTS WITH METASTATIC EPIDERMAL GROWTH FACTOR RECEPTOR (EGFR) T790M MUTATION-POSITIVE-NON-SMALL-CELL LUNG CANCER (NSCLC), AS DETECTED BY AN FDAAPPROVED TEST, WHO HAVE PROGRESSED ON OR AFTER EGFR TKI THERAPY	Converted	3/30/2017
DI A 704005	PRAXBIND	IDARUCIZUMAB	BOEHRINGER INGELHEIM	2/40/0045	10/46/0045	7.0	FOR THE TREATMENT OF PATIENTS TREATED WITH PRADAXA® WHEN REVERSAL OF THE ANTICOAGULANT EFFECTS OF DABIGATRAN IS NEEDED FOR EMERGENCY SURGERY/URGENT PROCEDURES AND IN LIFE-	Converted	4/42/2049
BLA 761025	PRAXBIND	IDAKUCIZUMAB	PHARMACEUTICALS	2/19/2015	10/16/2015	7.9	THREATENING OR UNCONTROLLED BLEEDING FOR THE TREATMENT OF PATIENTS WITH METASTATIC,	Converted	4/12/2018
BLA 125514 Supplement 5	KEYTRUDA	PEMBROLIZUMAB	MERCK SHARP & DOHME	4/2/2015	10/2/2015	6.0	PD-L1 POSITIVE, NON-SMALL CELL LUNG CANCER (NSCLC), AS DETERMINED BY AN FDA-APPROVED TEST, WITH DISEASE PROGRESSION ON OR AFTER PLATINUM-CONTAINING CHEMOTHERAPY	Converted	10/24/2016
BLA 125554 Supplement 2	OPDIVO	NIVOLUMAB	BRISTOL MYERS SQUIBB	3/30/2015	9/30/2015	6.0	IN COMBINATION WITH IPILIMUMAB, FOR THE TREATMENT OF PATIENTS WITH BRAF V600 WILD-TYPE, UNRESECTABLE OR METASTATIC MELANOMA	Converted	3/7/2019
2o Z	-: 20		1 1 2 3 1 3 2		2. 22.20.0	2.0	OTTITUDE OF TABLE OF THE TABLE TO THE TO THE TABLE OF THE	22/01.00	3/1/2018

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							FOR THE TREATMENT OF PATIENTS WITH TRANSFUSIONAL IRON OVERLOAD DUE TO		
							THALASSEMIA SYNDROMES WHEN CURRENT		
NDA 208030	FERRIPROX	DEFERIPRONE	APOPHARMA	11/17/2014	9/9/2015	9.7	CHELATION THERAPY IS INADEQUATE	Not Yet Converted	
							FOR THE TREATMENT OF CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS (TRANSFUSIONAL		
			NOVARTIS				HEMOSIDEROSIS) IN PATIENTS 2 YEARS OF AGE AND		
NDA 206910	JADENU	DEFERASIROX	PHARMACEUTICALS	5/30/2014	3/30/2015	10.0	OLDER	Converted	5/11/2018
							FOR THE TREATMENT OF CHRONIC IRON OVERLOAD IN		
							PATIENTS 10 YEARS OF AGE AND OLDER WITH NON- TRANSFUSION-DEPENDENT THALASSEMIA (NTDT)		
							SYNDROMES AND WITH A LIVER IRON CONCENTRATION		
							(LIC) OF AT LEAST 5 MILLIGRAMS OF IRON PER GRAM OF		
		BEEEB 101001	NOVARTIS	= 100 100 1	0/00/00/		LIVER DRY WEIGHT (MG FE/G DW) AND A SERUM		=/00/0000
NDA 206910	JADENU	DEFERASIROX	PHARMACEUTICALS	5/30/2014	3/30/2015	10.0	FERRITIN GREATER THAN 300 MCG/L	Converted	7/23/2020
							IN COMBINATION WITH BORTEZOMIB (BTZ) AND DEXAMETHASONE(DEX) FOR THE TREATMENT OF		
							PATIENTS WITH MULTIPLE MYELOMA (MM) WHO HAVE		
			NOVARTIS				RECEIVED AT LEAST 2 PRIOR REGIMENS, INCLUDING		
NDA 205353	FARYDAK	PANOBINOSTAT	PHARMACEUTICALS	3/24/2014	2/23/2015	11.0	BORTEZOMIB AND AN IMMUNOMODULATORY AGENT	Not Yet Converted	
							IN COMBINATION WITH LETROZOLE FOR THE TREATMENT OF POSTMENOPAUSAL WOMEN WITH		
							ESTROGEN RECEPTOR(ER)-POSITIVE, HUMAN		
							EPIDERMAL GROWTH FACTOR RECEPTOR 2 (HER2)-		
							NEGATIVE ADVANCED BREAST CANCER AS INITIAL		
NDA 207103	IBRANCE	PALBOCICLIB	PFIZER	8/13/2014	2/3/2015	5.7	ENDOCRINE-BASED THERAPY FOR THEIR METASTATIC DISEASE	Converted	3/31/2017
145/120/100	IDIOWOL	TALBOOICEIB	TTIZEN	0/10/2014	2/0/2010	0.7	FOR THE TREATMENT OF UNRESECTABLE OR	Converted	3/31/2017
							METASTATIC MELANOMA AND DISEASE PROGRESSION		
			BRISTOL MYERS				FOLLOWING IPILIMUMAB AND, IF BRAF V600 MUTATION		
BLA 125554	OPDIVO	NIVOLUMAB	SQUIBB	7/30/2014	12/22/2014	4.8	POSITIVE, A BRAF INHIBITOR	Converted	3/7/2019
							FOR PATIENTS WITH DELETERIOUS OR SUSPECTED DELETERIOUS GERMLINE BRCA MUTATED ADVANCED		
			ASTRAZENECA				OVARIAN CANCER WHO HAVE BEEN TREATED WITH		
NDA 206162	LYNPARZA	OLAPARIB	PHARMACEUTICALS	2/3/2014	12/19/2014	10.5	THREE OR MORE PRIOR LINES OF CHEMOTHERAPY	Converted	8/17/2017
							FOR THE TREATMENT OF PHILADELPHIA CHROMOSOME		
							NEGATIVE RELAPSED OR REFRACTORY B-CELL PRECURSOR ACUTE LYMPHOBLASTIC LEUKEMIA (ALL)		
BLA 125557	BLINCYTO	BLINATUMOMAB	AMGEN	9/19/2014	12/3/2014	2.5	The series (The series and the serie	Converted	7/11/2017
							FOR THE TREATMENT OF PATIENTS WITH		
			MERCK SHARP &				UNRESECTABLE OR METASTATIC MELANOMA AND DISEASE PROGRESSION FOLLOWING IPILIMUMAB AND,		
BLA 125514	KEYTRUDA	PEMBROLIZUMAB	DOHME	11/22/2013	9/4/2014	9.4	IF BRAF V600 MUTATION POSITIVE. A BRAF INHIBITOR	Converted	12/18/2015
	1						FOR THE TREATMENT OF RELAPSED FOLLICULAR B-		
							CELL NON-HODGKIN LYMPHOMA (FL) IN PATIENTS WHO		
							HAVE RECEIVED AT LEAST TWO PRIOR SYSTEMIC THERAPIES AND RELAPSED SMALL LYMPHOCYTIC		
							LYMPHOMA (SLL) IN PATIENTS WHO HAVE RECEIVED AT		
							LEAST TWO PRIOR SYSTEMIC THERAPIES		
NDA 205858	ZYDELIG	IDELALISIB	GILEAD SCIENCES	9/11/2013	7/23/2014	10.4		Not Yet Converted	
NDA 206256	BELEODAQ	BELINOSTAT	SPECTRUM PHARMACEUTICALS	12/9/2013	7/3/2014	6.8	FOR THE TREATMENT OF PATIENTS WITH RELAPSED OR REFRACTORY PERIPHERAL T-CELL LYMPHOMA	Not Yet Converted	
.10/1200200	SELECTION	SELINOOTAT		.2/3/2013	170/2017	0.0	FOR THE TREATMENTOF PATIENTS WITH ANAPLASTIC	Not 10t Convented	
							LYMPHOMA KINASE (ALK)-POSITIVE METASTATIC NON-		
							SMALL CELL LUNG CANCER (NSCLC) WHO HAVE		
NDA 205755	ZYKADIA	CERITINIB	NOVARTIS PHARMACEUTICALS	12/24/2013	4/29/2014	4.1	PROGRESSED ON OR ARE INTOLERANT TO CRIZOTINIB	Converted	5/26/2017
NDA 200730	ZINADIA	CERTINID	1 17 INWAGED HOAES	12/24/2013	4/20/2014	7.1	FOR THE TREATMENT OF ORTHOSTATIC DIZZINESS.	Converted	3/20/2017
	ĺ	1					LIGHTHEADEDNESS, OR THE "FEELING THAT YOU ARE		
	ĺ	1					ABOUT TO BLACK OUT" IN ADULT PATIENTS WITH		
	ĺ	1					SYMPTOMATIC NEUROGENIC ORTHOSTATIC		
							HYPOTENSION CAUSED BY PRIMARY AUTONOMIC FAILURE (PARKINSON'S DISEASE, MULTIPLE SYSTEM		
							ATROPHY, AND PURE AUTONOMIC FAILURE), DOPAMINE		
NID A GOOGE							BETA-HYDROXYLASE DEFICIENCY, AND NON-DIABETIC		
NDA 203202	NORTHERA	DROXIDOPA	LUNDBECK NA LTD	9/28/2011	2/18/2014	28.7	AUTONOMIC NEUROPATHY	Not Yet Converted	

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NDA 205552 Original 2	IMBRUVICA	IBRUTINIB	PHARMACYCLICS	6/28/2013	2/12/2014	7.5	FOR THE TREATMENT OF PATIENTS WITH CHRONIC LYMPHOCYTIC LEUKEMIA (CLL) WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY	Converted	7/28/2014
NDA 202806 Supplement 2	TAFINLAR	DABRAFENIB	NOVARTIS PHARMACEUTICALS	7/9/2013	1/9/2014	6.0	IN COMBINATION WITH TRAMETINIB, FOR THE TREATMENT OF PATIENTS WITH UNRESECTABLE OR METASTATIC MELANOMA WITH BRAF V600E OR V600K MUTATIONS, AS DETECTED BY AN FDA-APPROVED TEST	Converted	11/20/2015
NDA 204114 Supplement 1	MEKINIST	TRAMETINIB	NOVARTIS PHARMACEUTICALS	7/8/2013	1/8/2014	6.0	IN COMBINATION WITH DABRAFENIB, IS INDICATED FOR THE TREATMENT OF PATIENTS WITH UNRESECTABLE OR METASTATIC MELANOMA WITH BRAF V600E OR V600K MUTATIONS, AS DETECTED BY AN FDA-APPROVED TEST	Converted	11/20/2015
NDA 205552 Original 1	IMBRUVICA	IBRUTINIB	PHARMACYCLICS	6/28/2013	11/13/2013	4.5	FOR THE TREATMENT OF PATIENTS WITH MANTLE CELL LYMPHOMA (MCL)	Not Yet Converted	
BLA 125409 Supplement 51	PERJETA	PERTUZUMAB	GENENTECH	5/1/2013	9/30/2013	5.0	IN COMBINATION WITH TRASTUZUMAB AND DOCETAXEL FOR THE NEOADJUVANT TREATMENT OF PATIENTS WITH HER2-POSITIVE, LOCALLY ADVANCED, INFLAMMATORY, OR EARLY STAGE BREAST CANCER (EITHER GREATER THAN 2 CM IN DIAMETER OR NODE POSITIVE) AS PART OF A COMPLETE TREATMENT REGIMEN FOR EARLY BREAST CANCER	Converted	12/20/2017
BLA 125151 Supplement 184	ELAPRASE	IDURSULFASE	SHIRE HUMAN GENETIC THERAPIES	9/24/2012	6/24/2013	9.0	PROVIDES FOR ADDITIONAL SAFETY AND EFFICACY INFORMATION FOR THE TREATMENT OF PATIENTS WITH HUNTER SYNDROME 5 YEARS OF AGE AND YOUNGER	Not Yet Converted	
NDA 204026	POMALYST	POMALIDOMIDE	CELGENE	4/10/2012	2/8/2013	10.0	FOR THE TREATMENT OF PATIENTS WITH MULTIPLE MYELOMA WHO HAVE RECEIVED AT LEAST TWO PRIOR THERAPIES INCLUDING LENALIDOMIDE AND BORTEZOMIB AND HAVE DEMONSTRATED DISEASE PROGRESSION ON OR WITHIN 60 DAYS OF COMPLETION OF THE LAST THERAPY	Converted	4/23/2015
NDA 021882 Supplement 15	EXJADE	DEFERASIROX	NOVARTIS PHARMACEUTICALS	12/23/2011	1/23/2013	13.1	FOR THE TREATMENT OF CHRONIC IRON OVERLOAD IN PATIENTS 10 YEARS OF AGE AND OLDER WITH NON-TRANSFUSION DEPENDENT THALASSEMIA (NTDT) SYNDROMES AND WITH A LIVER IRON CONCENTRATION (LIC) OF AT LEAST5 MILLIGRAMS OF IRON PER GRAM OF LIVER DRY WEIGHT (MG FE/G DW) AND A SERUM FERRITIN GREATER THAN 300 MCG/L	Converted	7/23/2020
NDA 204384	SIRTURO	BEDAQUILINE	JANSSEN RESEARCH AND DEVELOPMENT	6/29/2012	12/28/2012	6.0	FOR THE TREATMENT OF, AS COMBINATION THERAPY, ADULTS (2: 18 YEARS) WITH PULMONARY MULTI-DRUG RESISTANT TUBERCULOSIS (MDR-TB)	Not Yet Converted	772072020
NDA 203469	ICLUSIG	PONATINIB	ARIAD PHARMACEUTICALS	9/27/2012	12/14/2012	2.6	FOR THE TREATMENT OF ADULT PATIENTS WITH CHRONIC PHASE, ACCELERATED PHASE, OR BLAST PHASE CHRONIC MYELOID LEUKEMIA (CML) THAT IS RESISTANT OR INTOLERANT TO PRIOR TYROSINE KINASE INHIBITOR THERAPY OR PHILADELPHIA CHROMOSOME POSITIVE ACUTE LYMPHOBLASTIC LEUKEMIA (PH-ALL) THAT IS RESISTANT OR INTOLERANT TO PRIOR TYROSINE KINASE INHIBITOR THERAPY	Converted	11/28/2016
NDA 203585	SYNRIBO	OMACETAXINE MEPESUCCINATE	TEVA PHARMACEUTICALS INTERNATIONAL	3/30/2012	10/26/2012	6.9	FOR THE TREATMENT OF ADULT PATIENTS WITH CHRONIC OR ACCELERATED PHASE CHRONIC MYELOID LEUKEMIA (CML) WITH RESISTANCE AND/OR INTOLERANCE TO TWO OR MORE TYROSINE KINASE INHIBITORS (TKI)	Converted	2/10/2014
NDA 203985	AFINITOR DISPERZ	EVEROLIMUS	NOVARTIS PHARMACEUTICALS	2/29/2012	8/29/2012	6.0	FOR THE TREATMENT OF PEDIATRIC AND ADULT PATIENTS WITH TUBEROUS SCLEROSIS COMPLEX (TSC) FOR THE TREATMENT OF SUBEPENDYMAL GAIANT CELL ASTROCYTOMA (SEGA) THAT REQUIRES THERAPEUTIC INTERVENTION BUT CANNOT BE CURATIVELY RESECTED	Converted	1/29/2016
	2101 2112			-, -0, -0, 12	3/20/2012	0.0	NEOLOTED	Conveneu	1/29/2010

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NDA 202497	MARQIBO	VINCRISTINE SULFATE (LIPOSOMAL)	TALON THERAPEUTICS	7/12/2011	8/9/2012	13.0	FOR THE TREATMENT OF ADULTS WITH PHILADELPHIA (PH) CHROMOSOME NEGATIVE (-) ACUTE LYMPHOBLASTIC LEUKEMIA (ALL) IN SECOND RELAPSE OR GREATER RELAPSE OR WHOSE DISEASE HAS PROGRESSED FOLLOWING TWO OR GREATHER TREATMENT LINES OF ANTI-LEUKEMIA THERAPIES	Not Yet Converted	
NDA 202714	KYPROLIS	CARFILZOMIB	ONYX THERAPEUTICS	9/27/2011	7/20/2012	9.8	FOR THE TREATMENT OF PATIENTS WITH MULTIPLE MYELOMA WHO HAVE RECEIVED AT LEAST TWO PRIOR THERAPIES INCLUDING BORTEZOMIB AND AN IMMUNOMODULATORY AGENT AND HAVE DEMONSTRATED DISEASE PROGRESSION ON OR WITHIN 60 DAYS OF COMPLETION OF THE LAST THERAPY	Converted	1/21/2016
NDA 022334 Supplement 17	AFINITOR	EVEROLIMUS	NOVARTIS PHARMACEUTICALS	12/19/2011	4/26/2012	4.2	TREATMENT OF ADULTS WITH RENAL ANGIOMYOLIPOMA AND TUBEROUS SCLEROSIS COMPLEX (TSC) NOT REQUIRING IMMEDIATE SURGERY	Converted	2/18/2016
NDA 021825	FERRIPROX	DEFERIPRONE	APOPHARMA	1/30/2009	10/14/2011	32.4	FOR THE TREATMENT OF TRANSFUSIONAL IRON OVERLOAD DUE TO THALASSEMIA SYNDROMES WHEN CURRENT CHELATION THERAPY IS INADEQUATE	Not Yet Converted	
NDA 202570	XALKORI	CRIZOTINIB	PF PRISM CV	3/30/2011	8/26/2011	4.9	FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC NON-SMALL CELL LUNG CANCER (NSCLC) THAT IS ANAPLASTIC LYMPHOMA KINASE (ALK)-POSITIVE AS DETECTED BY AN FDA APPROVED TEST	Converted	11/20/2013
BLA 125388	ADCEDTRIS	BRENTUXIMAB VEDOTIN	SEATTLE GENETICS	2/28/2011	8/19/2011	5.7	FOR THE TREATMENT OF PATIENTS WITH HODGKIN LYMPHOMA AFTER FAILURE OF AUTOLOGOUS STEM CELL TRANSPLANT (ASCT) OR AFTER FAILURE OF AT LEAST TWO PRIOR MULTI-AGENT CHEMOTHERAPY REGIMENS IN PATIENTS WHO ARE NOT ASCT CANDIDATES	Converted	8/17/2015
BLA 125388 Supplement 6	ADCETRIS	BRENTUXIMAB VEDOTIN	SEATTLE GENETICS	2/28/2011	8/19/2011	5.7	FOR THE TREATMENT OF PATIENTS WITH SYSTEMIC ANAPLASTIC LARGE CELL LYMPHOMA (SALCL) AFTER FAILURE OF AT LEAST ONE PRIOR MULIT-AGENT CHEMOTHERPAY REGIMEN	Converted	3/20/2018
NDA 022393 Supplement 4	ISTODAX	ROMIDEPSIN	CELGENE	12/17/2010	6/16/2011	6.0	THE TREATMENT OF PERIPHERAL T-CELL LYMPHOMA (PTCL) IN PATIENTS WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY	Not Yet Converted	
NDA 021945	MAKENA	HYDROXYPROGES TERONE CAPROATE	AMAG PHARMA USA	4/20/2006	2/3/2011	57.5	TO REDUCE THE RISK OF PRETERM BIRTH IN WOMEN WITH A SINGLETON PREGNANCY WHO HAVE A HISTORY OF SINGLETON SPONTANEOUS PRETERM BIRTH	Not Yet Converted	
NDA 022334 Supplement 6	AFINITOR	EVEROLIMUS	NOVARTIS PHARMACEUTICALS	4/30/2010	10/29/2010	6.0	FOR THE TREATMENT OF SUBEPENDYMAL GIANT CELL ASTROCYTOMA (SEGA) ASSOCIATED WITH TUBEROUS SCLEROSIS (TS) WHO REQUIRE THERAPEUTIC INTERVENTION BUT ARE NOT CANDIDATES FOR CURATIVE SURGICAL RESECTION	Converted	1/29/2016
NDA 021986 Supplement 8	SPRYCEL	DASATINIB	BRISTOL MYERS SQUIBB	4/28/2010	10/28/2010	6.0	FOR THE TREATMENT OF NEWLY DIAGNOSED ADULTS WITH PHILADELPHIA CHROMOSOME-POSITIVE (PH+) CML IN CHRONIC PHASE	Converted	8/12/2015
NDA 022068 Supplement 5	TASIGNA	NILOTINIB	NOVARTIS PHARMACEUTICALS	12/21/2009	6/17/2010	5.9	FOR THE TREATMENT OF NEWLY DIAGNOSED ADULT PATIENTS WITH PHILADELPHIA CHROMOSOME POSITIVE CHRONIC MYELOID LEUKEMIA IN CHRONIC PHASE	Converted	1/27/2015
NDA 022059 Supplement 7	TYKERB	LAPATINIB	NOVARTIS PHARMACEUTICALS	3/31/2009	1/29/2010	10.0	TYKERB IN COMBINATION WITH LETROZOLE IS FOR THE TREATMENT OF POST MENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE METASTATIC BREAST CANCER THAT OVEREXPRESS THE HER2 RECEPTOR FOR WHOM HORMONAL THERAPY IS INDICATED	Converted	12/6/2018
BLA 125326	ARZERRA	OFATUMUMAB	NOVARTIS PHARMACEUTICALS	1/30/2009	10/26/2009	8.8	FOR THE TREATMENT OF PATIENTS WITH CHRONIC LYMPHOCYTIC LEUKEMIA (CLL) REFRACTORY TO FLUDARABINE AND ALEMTUZUMAB	Converted	4/17/2014
NDA 022468	FOLOTYN	PRALATREXATE	ALLOS THERAPEUTICS	3/24/2009	9/24/2009	6.0	FOR THE TREATMENT OF RELAPSED OR REFRACTORY PERIPERAL T-CELL LYMPHOMA	Not Yet Converted	
BLA 125085 Supplement 169	AVASTIN	BEVACIZUMAB	GENENTECH	11/3/2008	5/5/2009	6.0	FOR THE TREATMENT OF GLIOBLASTOMA WITH PROGRESSIVE DISEASE FOLLOWING PRIOR THERAPY	Converted	12/5/2017

NDA 021588		IMATINIB	NOVARTIS				FOR THE ADJUVANT TREATMENT OF ADULT PATIENTS FOLLOWING COMPLETE GROSS RESECTION OF KIT (CD117) POSITIVE GASTROINTESTINAL STROMAL		
Supplement 25	GLEEVEC	MESYLATE	PHARMACEUTICALS	6/24/2008	12/19/2008	5.9	TUMORS (GIST)	Converted	1/31/2012
		FLUDARABINE					FOR THE TREATMENT OF ADULTS PTS WITH B CELL CHRONIC LYMPHOCYTIC LEUEMIA (CLL) WHOSE DISEASE HAS NOT RESPONDED TO OR WHO HAVE NOT RESPONDED TO OR HAS PROGRESSED DURING OR AFTER TREATMENT WITH AT LEAST ONE STANDARD ALKYLATING-AGENT CONTAINING REGIMEN		
NDA 022273	OFORTA	PHOSPHATE	SANOFI AVENTIS	11/19/2007	12/18/2008	13.0	ALKIEKTING AGENT GONTAINING REGIMEN	Not Converted-Application Withdrawn	12/31/2011
NDA 022291	PROMACTA	ELTROMBOPAG	NOVARTIS PHARMACEUTICALS	12/19/2007	11/20/2008	11.1	FOR THE TREATMENT THROMBOCYTOPNEIA IN PATIENTS WITH CHRONIC IMMUNE (DIDOPATHIC) THROMBOCYTOPENIA (ITP) WHO HAVE HAD AN INSUFFICIENT RESPONSE TO CORTICOSTEROIDS, IMMUNOGLOBULINS, OR SPLENECTOMY	Converted	2/25/2011
NDA 021462 Supplement 15	ALIMTA	PEMETREXED DISODIUM	ELI LILLY AND COMPANY	8/28/2007	9/26/2008	13.0	FOR THE TREATMENT OF • NONSQUAMOUS NON-SMALL CELL LUNG CANCER: INITIAL TREATMENT IN COMBINATION WITH CISPLATIN. (1.1) • NONSQUAMOUS NON-SMALL CELL LUNG CANCER AS A SINGLE-AGENT AFTER PRIOR CHEMOTHERAPY (1.2)	Converted	7/2/2009
NDA 020634			JANSSEN	0.20.200	0,20,200		TO REDUCE THE INCIDENCE OR PROGRESSION OF DISEASE FOLLOWING EXPOSURE TO AEROSOLIZED BACILLUS ANTHRACIS IN PEDIATRIC PATIENTS (>6		
Supplement 47	LEVAQUIN	LEVOFLOXACIN	PHARMACEUTICALS	7/5/2007	5/5/2008	10.0	MONTH OF AGE AND OLDER)	Not Yet Converted	
NDA 020635 Supplement 51	LEVAQUIN	LEVOFLOXACIN	JANSSEN PHARMACEUTICALS	7/5/2007	5/5/2008	10.0	TO REDUCE THE INCIDENCE OR PROGRESSION OF DISEASE FOLLOWING EXPOSURE TO AEROSOLIZED BACILLUS ANTHRACIS IN PEDIATRIC PATIENTS (>6 MONTH OF AGE AND OLDER)	Not Converted-Application Withdrawn	7/21/2017
NDA 021721 Supplement 15	LEVAQUIN (oral solution)	LEVOFLOXACIN	JANSSEN PHARMACEUTICALS	7/5/2007	5/5/2008	10.0	TO REDUCE THE INCIDENCE OR PROGRESSION OF DISEASE FOLLOWING EXPOSURE TO AEROSOLIZED BACILLUS ANTHRACIS IN PEDIATRIC PATIENTS (>6 MONTH OF AGE AND OLDER)	Not Converted-Application Withdrawn	7/21/2017
BLA 125085 Supplement 91	AVASTIN	BEVACIZUMAB	GENENTECH	5/24/2006	2/22/2008	21.0	NEW INDICATION FOR USE IN COMBINATION WITH PACLITAXEL FOR THE TREATMENT OF PATIENTS WHO HAVE NOT RECEIVED CHEMOTHERAPY FOR METASTATIC HER2 NEGATIVE BREAST CANCER	Not Converted-Indication Withdrawn	11/18/2011
NDA 022187	INTELENCE	ETRAVIRINE	JANSSEN PRODUCTS LP	7/18/2007	1/18/2008	6.0	IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS FOR THE TREATMENT OF HIV-1 INFECTION IN TREATMENT-EXPERIENCED ADULT PATIENTS WHO HAVE EVIDENCE OF VIRAL REPLICATIONS AND HIV-1 STRAINS RESISTANT TO NON-NUCLELOSIDE REVERSE TRANSCRIPTASE INHIBITOR (NNRTI) AND OTHER ANTIRETROVIRAL AGENT	Converted	11/24/2009
NDA 022068	TASIGNA	NILOTINIB	NOVARTIS PHARMACEUTICALS	9/29/2006	10/29/2007	13.0	FOR THE TREATMENT OF CHRONIC PHASE (CP) AND ACCELERATED PHASE (AC) PHILADELPHIA CHROMOSONE POSITIVE CHRONIC MYELOGENOUS LEUKEMIA(CML)IN ADULT PATIENTS RESISTANT TO OR INTOLERANT TO PRIOR THERAPY THAT INCLUDED GLEEVEC (IMATINIB)	Converted	1/14/2011
			MERCK SHARP &				IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS, FOR THE TREATMENT OF HIV 1 INFECTION IN TREATMENT-EXPERIENCED ADULT PATIENTS WHO HAVE EVIDENCE OF VIRAL REPLICATION AND HIV-1 STRAINS RESISTANT TO MULTIPLE ANTIRETORVIRAL		
NDA 022145	ISENTRESS	RALTEGRAVIR	DOHME	4/13/2007	10/12/2007	6.0	AGENTS FOR THE TREATMENT OF PATIENTS WITH CCR5-TROPIC	Converted	1/29/2009
NDA 022128	SELZENTRY	MARAVIROC	VIIV HEALTHCARE	12/20/2006	8/6/2007	7.5	HIV-1	Converted	11/25/2008
NDA 021588 Supplement 16	GLEEVEC	IMATINIB MESYLATE	NOVARTIS PHARMACEUTICALS	3/28/2006	9/27/2006	6.0	FOR THE TREATMENT OF NEWLY DIAGNOSED PHILADELPHIA POSITIVE CML IN PEDIATRIC PATIENTS	Converted	4/1/2011
BLA 125147	VECTIBIX	PANITUMUMAB	AMGEN	3/29/2006	9/27/2006	6.0	FOR THE TREATMENT OF EGFR-EXPRESSING, METASTATIC COLORECTAL CARCINOMA WITH DISEASE PROGRESSION ON OR FOLLOWING FLUOROPYRIMIDINE-, OXALIPLATIN-, AND IRINOTECAN-CONTAINING CHEMOTHERAPY REGIMENS	Commented	5/22/2044
JEN 120141	VECTION	a a a a a a a a a a a a a a a a a a	/ WIOLIY	3/23/2000	3/2//2000	0.0	OFFICIAL LART REGIMENS	Converted	5/23/2014

NDA 021986 SPRYCEL DASATINIB BRISTOL MYERS SQUIBB 12/28/2005 6/28/2006 6.0 FOR THE TREATMENT OF CHRONIC MYELOID LEUKEMIA WITH RESISTANCE OR INTOLERANCE TO PRIOR THE TREATMENT OF CHRONIC MYELOID LEUKEMIA WITH RESISTANCE OR INTOLERANCE TO PRIOR THE REATMENT OF CHIMAN IMMUNODEFICIENCY VIRUS (HIV) INFECTION IN ANTIRETROVIRAL TREATMENT EXPERIENCED ADULT PATIENTS, SUCH AS THOSE WITH HIV-1 STRAINS RESISTANT TO MORE THAN ONE PRODUCTS LP 12/23/2005 6/23/2006 6.0 PROTESSE INHIBITOR Converted PROTECTION IN ANTIRETROVIRAL TREATMENT OF PROTESSE INHIBITOR CONVERTED TO PROTESSE INHI	5/21/2009 10/21/2008 6/19/2014 2/2/2007 4/30/2010 5/11/2018
NDA 021976 PREZISTA DARUNAVIR PRODUCTS LP 12/23/2005 6/23/2006 6.0 PROTEASE INHIBITOR CONVERTED ADULT PATIENTS, SUCH AS THOSE WITH HIV-1 STRAINS RESISTANT TO MORE THAN ONE PROTEASE INHIBITOR CONVERTED ADULT PATIENTS, SUCH AS THOSE WITH HIV-1 STRAINS RESISTANT TO MORE THAN ONE PROTEASE INHIBITOR CONVERTED ADULT PATIENTS WITH NEWLY DIAGNOSED MULTIPLE MYELOMA CONVERTED ADULT PATIENTS WITH THE PROTECTION OF PATIENTS WITH THE PROTECTION OF PATIENTS WITH THE PROTECTION OF PATIENTS WITH THE PATIENT OF PATIENTS WITH THE	6/19/2014 2/2/2007 4/30/2010 5/11/2018
NDA 021976 PREZISTA DARUNAVIR PRODUCTS LP 12/23/2005 6/23/2006 6.0 PROTEASE INHIBITOR Converted NDA 021430 Original 1 THALOMID THALIDOMIDE CELGENE 12/23/2003 5/25/2006 29.1 DIAGNOSED MULTIPLE MYELOMA CONVERTED NDA 021968 Original 1 SUTENT SUTENT MALEATE SUNITINIB MALEATE SUNITINIB NOVARTIS Supplement 12 FEMARA LETROZOLE PHARMACEUTICALS SUPPLEMENTOR SUNITINIB NOVARTIS	6/19/2014 2/2/2007 4/30/2010 5/11/2018
Original 1 THALOMID THALIDOMIDE CELGENE 12/23/2003 5/25/2006 29.1 DIAGNOSED MULTIPLE MYELOMA COnverted ORIGINAL SUNTINIB SUNTINIB SUNTENT MALEATE SUNTINIB MALEATE SUNTENATIONAL 8/11/2005 1/26/2006 5.5 FOR THE TREATMENT OF ADVANCED RENAL CELL CARCINOMA CONVERTED CONVERTED SUNTINIB MALEATE SUNTENATIONAL 8/11/2005 1/26/2006 5.5 FOR THE TREATMENT OF ADVANCED RENAL CELL CARCINOMA CONVERTED CONVERTED SUNTINIB MALEATE SUNTENATIONAL 8/11/2005 1/26/2006 5.5 FOR THE ADJUVANT TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER CONVERTED SUNTENATION OF A S	2/2/2007 4/30/2010 5/11/2018
NDA 021968 Original 1 SUTENT NDA 020726 Supplement 12 FEMARA LETROZOLE NOVARTIS NDA 021882 EXJADE DEFERASIROX NOVARTIS NDA 021877 ARRANON NELARABINE SUNITINIB MALEATE PHARMACEUTICALS 8/11/2005 1/26/2006 8/11/2005 1/26/2006 1/26/2006 1/26/2006 1/26/2006 1/26/2006 5.5 FOR THE ADJUVANT TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER FOR THE TREATMENT OF CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER FOR THE TREATMENT OF CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER Converted FOR THE TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER Converted FOR THE TREATMENT OF PATIENTS WITH T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUK	4/30/2010 5/11/2018
Original 1 SUTENT MALEATE INTERNATIONAL 8/11/2005 1/26/2006 5.5 Converted NDA 020726 Supplement 12 FEMARA LETROZOLE NOVARTIS PHARMACEUTICALS 6/28/2005 12/28/2005 6.0 BREAST CANCER NOVARTIS NDA 021882 EXJADE DEFERASIROX PHARMACEUTICALS 5/2/2005 11/2/2005 6.0 G.0 DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER FOR THE TREATMENT OF CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER Converted FOR THE TREATMENT OF PATIENTS WITH T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSEED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS CONVERTED CONVERTE	4/30/2010 5/11/2018
NDA 020726 Supplement 12 FEMARA LETROZOLE PHARMACEUTICALS 6/28/2005 12/28/2005 6.0 WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER FOR THE TREATMENT OF CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER Converted FOR THE TREATMENT OF PATIENTS 2 YEARS OF AGE AND OLDER CONVERTED CONVERTED THYPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS CONVERTED CON	5/11/2018
NDA 021882 EXJADE DEFERASIROX PHARMACEUTICALS 5/2/2005 11/2/2005 6.0 DUE TO BLOOD TRANSFUSIONS IN PATIENTS 2 YEARS OF AGE AND OLDER CONVerted FOR THE TREATMENT OF PATIENTS WITH T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS NDA 021877 ARRANON NELARABINE GLAXOSMITHKLINE 4/29/2005 10/28/2005 6.0 REGIMENS CONVERTED CO-ADMINISTERED WITH 200 MG OF RITONAVIR FOR	
LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY NDA 021877 ARRANON NELARABINE GLAXOSMITHKLINE 4/29/2005 10/28/2005 6.0 REGIMENS CO-ADMINISTERED WITH 200 MG OF RITONAVIR FOR	
CO-ADMINISTERED WITH 200 MG OF RITONAVIR FOR	
	7/31/2019
BOEHRINGER INFECTED ADULT PATIENTS WITH EVIDENCE OF VIRAL REPLICATION, WHO ARE HIGHLY TREATMENT-INGELHEIM EXPERIENCED OR HAVE HIV-1 STRAINS RESISTANT TO	
NDA 021814 APTIVUS TIPRANAVIR PHARMACEUTICALS 12/22/2004 6/22/2005 6.0 MULTIPLE PROTEASE INHIBITORS Converted	10/4/2007
FOR THE TREATMENT OF PEDIATRIC PATIENTS 1 TO 21 YEARS OLD WITH RELAPSED OR REFRACTORY ACUTE LYMPHOCYTIC LEUKEMIA AFTER AT LEAST TWO PRIOR	
NDA 021673 CLOLAR CLOFARABINE GENZYME 3/30/2004 12/28/2004 9.0 REGIMENS Not Yet Convert	ied
EXPAND THE INDICATION TO INCLUDE PATIENTS WITH RELAPSED OR REFRACTORY, LOW GRADE, FOLLICULAR TOSITUMOMAB BLA 125011 AND IODINE I 131 EXPAND THE INDICATION TO INCLUDE PATIENTS WITH RELAPSED OR REFRACTORY, LOW GRADE, FOLLICULAR OR TRANSFORMED CD20 POSITIVE NON-HODGKIN'S LYMPHOMA WHO HAVE NOT RECEIVED RITUXIMAB	
Supplement 24 BEXXAR TOSITUMOMAB GLAXOSMITHKLINE 7/3/2004 12/22/2004 5.7 Not Converted-Indication NDA 020634 JANSSEN FOR THE TREATMENT OF INHALATIONAL ANTHRAX	n Withdrawn 10/23/2013
Supplement 35 LEVAQUIN LEVOFLOXACIN PHARMACEUTICALS 5/26/2004 11/24/2004 6.0 (POST-EXPOSURE) Not Yet Convert	ied
NDA 020635 Supplement 35 LEVAQUIN LEVOFLOXACIN PHARMACEUTICALS 5/26/2004 11/24/2004 6.0 (POST-EXPOSURE) FOR THE TREATMENT OF INHALATIONAL ANTHRAX (POST-EXPOSURE) Not Converted-Application	n Withdrawn 7/21/2017
NDA 021721 JANSSEN FOR THE TREATMENT OF INHALATIONAL ANTHRAX Supplement 3 AQUIN (oral solu LEVOFLOXACIN PHARMACEUTICALS 11/12/2004 11/24/2004 0.4 (POST-EXPOSURE) Not Converted-Application	n Withdrawn 7/21/2017
PROVIDES FOR ADDING THE INFUSION OF REMODULIN (TREPROSTINIL SODIUM) 1, 2.5, 5 & 10 MG/ML INJECTION VIA AN INDWELLING CENTRAL VENOUS CATHETER TO NDA 021272 Supplement 2 REMODULIN SODIUM THERAPEUTICS 1/30/2004 11/24/2004 9.8 ARTERIAL HYPERTENSION (PAH) Converted	3/20/2006
Supplement 2 REMODULIN SODIUM THERAPEUTICS 1/30/2004 11/24/2004 9.8 ARTERIAL HYPERTENSION (PAH) Converted FOR THE TREATMENT OF PATIENTS WITH RELAPSING	3/20/2006
BLA 125104 TYSABRI NATALIZUMAB BIOGEN 5/24/2004 11/23/2004 6.0 FORMS OF MULTIPLE SCLEROSIS (MS) TO REDUCE THE FREQUENCY OF CLINICAL EXACERBATIONS Converted	6/5/2006
FOR THE EXTENDED ADJUVANT TREATMENT OF EARLY BREAST CANCER IN POSTMENOPAUSAL WOMEN WHO NOVARTIS Supplement 11 FEMARA LETROZOLE PHARMACEUTICALS 4/29/2004 10/29/2004 6.0 THERAPY Converted	4/30/2010
CONCOMITANTLY ADMINISTERED WITH GONAL-F	4/30/2010
(FOLLITROPIN ALFA FOR INJECTION) FOR STIMULATION OF FOLLICULAR DEVELOPMENT IN INFERTILE HYPOGONADOTROPIC HYPOGONADAL WOMEN WITH	
NDA 021322 LUVERIS LUTROPIN ALPHA EMD SERONO 5/1/2001 10/8/2004 41.3 PROFOUND LH DEFICIENCY(LH < 1.2 IU/L) Not Converted-Application	n Withdrawn 4/12/2016

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NDA 021677		PEMETREXED	ELI LILLY AND				AS A SINGLE AGENT FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATAIC NON- SMALL CELL LUNG CANCER AFTER PRIOR		
Original 1	ALIMTA	DISODIUM	COMPANY	11/4/2003	8/19/2004	9.5	CHEMOTHERPY IN COMBINATION WITH OTHER ANTIRETROVIRAL	Converted	7/2/2009
		DISOPROXIL					AGENTS FOR THE TREATMENT OF HIV INFECTION IN		
NDA 021752	TRUVADA	FUMARATE; EMTRICITABINE	GILEAD SCIENCES	3/12/2004	8/2/2004	4.7	ADULTS	Converted	3/8/2006
							IN COMBINATION WITH IRINOTECAN FOR THE TREATMENT OF EGFR-EXPRESSING METASTATIC		
			ELI LILLY AND				COLORECTAL CARCINOMA IN PATIENTS WHO ARE REFRACTORY TO IRINOTECAN-BASED CHEMOTHERAPY		
BLA 125084	CETUXIMAB	ERBITUX	COMPANY	8/14/2003	2/12/2004	6.0		Converted	7/6/2012
							AS A SINGLE AGENT FOR THE TREATMENT OF EGFR- EXPRESSING, METASTATIC COLORECTAL CARCINOMA		
BLA 125084	CETUXIMAB	ERBITUX	ELI LILLY AND COMPANY	8/14/2003	2/12/2004	6.0	IN PATIENTS WHO ARE INTOLERANT TO IRINOTECAN- BASED CHEMOTHERAPY	Converted	10/2/2017
							FOR THE TREATMENT OF PEDIATRIC PATIENTS WITH		
NDA 021335	OLEEVEO	IMATINIB	NOVARTIS	0/00/0000	F/00/0000	40.7	PH+ CHRONIC PHASE CML WHOSE DISEASE HAS RECURRED AFTER STEM CELL TRANSPLANT OR WHO		
Supplement 3	GLEEVEC	MESYLATE	PHARMACEUTICALS	6/28/2002	5/20/2003	10.7	ARE RESISTANT TO INTERFERON ALPHA THERAPY FOR THE TREATMENT OF MULTIPLE MYELOMA	Converted	9/27/2006
			MILLENNIUM				PATIENTS WHO HAVE RECEIVED AT LEAST TWO PRIOR THERAPIES AND HAVE DEMONSTRATED DISEASE		
NDA 021602	VELCADE	BORTEZOMIB	PHARMACEUTICALS	1/21/2003	5/13/2003	3.7	PROGRESSION ON THE LAST THERAPY	Converted	3/25/2005
							AS A MONOTHERAPY FOR THE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC		
			ASTRAZENECA UK				NON-SMALL CELL LUNG CANCER AFTER FAILURE OF BOTH PLATINUM-BASED AND DOCETAXEL		
NDA 021399	IRESSA	GEFITINIB	LTD	8/5/2002	5/5/2003	9.0	CHEMOTHERAPY	Not Converted-Application Withdrawn	4/25/2012
BLA 103979	FABRAZYME	AGALSIDASE BETA	GENZYME	6/23/2000	4/24/2003	34.0	FOR THE TREATMENT OF FABRY DISEASE	Not Yet Converted	
							IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS FOR THE TREATMENT OF HIV-1 INFECTION IN		
							TREATMENT EXPERIENCED PATIENTS WITH EVIDENCE OF HIV-1 REPLICATION DESPITE ONGOING		
NDA 021481	FUZEON	ENFUVIRTIDE	HOFFMAN LA ROCHE	9/16/2002	3/13/2003	5.9	ANTIRETROVIRAL THERAPY	Converted	10/15/2004
NDA 021335		IMATINIB	NOVARTIS				FOR THE TREATMENT OF NEWLY DIAGNOSED ADULT PATIENTS WITH PHILADELPHIA CHROMOSOME POSITIVE		
Supplement 4	GLEEVEC	MESYLATE	PHARMACEUTICALS	6/28/2002	12/20/2002	5.8	CHRONIC MYELOID LEUKEMIA (CML) FOR THE ADJUVANT TREATMENT OF POSTMENOPAUSAL	Converted	5/27/2009
NDA 020541 Supplement 10	ARIMIDEX	ANASTROZOLE	ANI PHARMACEUTICALS	3/5/2002	9/5/2002	6.0	WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY	Converted	9/16/2005
Заррієпієть то	AKIIVIIDEX	ANASTROZOLL	THANNACEOTICALS	3/3/2002	3/3/2002	0.0	BREAST CANCER IN COMBINATION WITH INFUSIONAL 5-FU/LV FOR THE	Converted	9/16/2005
							TREATMENT FOR PATIENTS WITH METASTATIC CARCINOMA OF THE COLON OR RECTUM WHOSE		
							DISEASE HAS PROGRESSED DURING OR WITHIN 6 MONTHS OF COMPLETION OF FIRST LINE THERAPY		
NDA 021492	ELOXATIN	OXALIPLATIN	SANOFI AVENTIS	6/24/2002	8/9/2002	4.5	WITH THE COMBINATION OF BOLUS 5 -FU/LV AND	Commission	4 10 10 0 0 4
		TREPROSTINIL	UNITED			1.5	IRINOTECAN TREATMENT OF PULMONARY ARTERIAL HYPERTENSION	Converted	1/9/2004
NDA 021272	REMODULIN	SODIUM	THERAPEUTICS	10/16/2000	5/21/2002	19.1	FOR THE TREATMENT OF PATIENTS WITH RELAPSED OR	Converted	3/20/2006
							REFRACTORY LOW-GRADE, FOLLICULAR, OR		
		IBRITUMOMAB	SPECTRUM				TRANSFORMED B-CELL NON-HODGKIN'S LYMPHOMA(NHL) OTHER THAN THOSE PATIENTS WITH		
BLA 125019	ZEVALIN	TIUXETAN	PHARMACEUTICALS	11/1/2000	2/19/2002	15.6	RITUXIMAB REFRACTORY FOLLICULAR NHL FOR THE TREATMENT OF PATIENTS WITH KIT (CD117)	Converted	9/3/2009
NDA 021335		IMATINIB	NOVARTIS				POSITIVE UNRESECTABLE AND/OR METASTATIC MALIGNANT GASTROINTESTINAL		
Supplement 1	GLEEVEC	MESYLATE	PHARMACEUTICALS	10/16/2001	2/1/2002	3.6	STROMAL TUMORS (GIST)	Converted	9/26/2008
		TENOFOVIR DISOPROXIL					FOR THE TREATMENT OF HIV-1 INFECTION IN ADULTS	·	
NDA 021356	VIREAD	FUMARATE	GILEAD SCIENCES	5/1/2001	10/26/2001	5.9		Converted	3/8/2006

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NDA 021335	GLEEVEC	IMATINIB MESYLATE	NOVARTIS PHARMACEUTICALS	2/27/2001	5/10/2001	2.4	FOR THE TREATMENT OF PATIENTS WITH CHRONIC MYELOID LEUKEMIA(CML) IN BLAST CRISIS, ACCELERATED PHASE, OR IN CHRONIC PHASE AFTER	Converted	40/0/000
NDA 021333	GLEEVEC	WESTLATE	FHARMACEUTICALS	2/21/2001	5/10/2001	2.4	FAILURE OF INTERFERON-ALPHA THERAPY FOR THE TREATMENT OF B-CELL CHRONIC LYMPHOCYTIC LEUKEMIA (B-CLL) IN PATIENTS WHO	Convened	12/8/2003
BLA 103948	CAMPATH	ALEMTUZUMAB	GEMZYME	12/23/1999	5/7/2001	16.5	HAVE BEEN TREATED WITH ALKYLATING AGENTS AND WHO HAVE FAILED FLUDARABINE THERAPY	Converted	9/19/2007
		ABACAVIR SULFATE;	-				ALONE OR IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS FOR THE TREATMENT OF HIV-		3/10/2001
NDA 021205	TRIZIVIR	LAMIVUDINE; ZIDOVUDINE	VIIV HEALTHCARE	12/17/1999	11/14/2000	10.9	1 INFECTION	Converted	5/13/2005
							IN COMBINATION WITH OTHER ANTIRETROVIRAL		
	KALETRA	LOPINAVIR:					AGENTS FOR THE TREATMENT OF HIV-1 INFECTION IN ADULTS AND PEDIATRIC PATIENTS AGE SIX MONTHS		
NDA 021226	(capsules)	RITONAVIR	ABBVIE	6/1/2000	9/15/2000	3.5	OR OLDER	Converted	11/27/2002
NDA 021251	KALETRA (oral solution)	LOPINAVIR: RITONAVIR	ABBVIE	6/1/2000	9/15/2000	3.5	IIN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS FOR THE TREATMENT OF HIV-1 INFECTION IN ADULTS AND PEDIATRIC PATIENTS AGE SIX MONTHS OR OLDER	Converted	11/27/2002
NDA 019537			BAYER HEALTHCARE				FOR THE TREATMENT OF INHALATIONAL ANTHRAX (POST-EXPOSURE)		
Supplement 38	CIPRO	CIPROFLOXACIN	PHARMACEUTICALS	3/1/2000	8/30/2000	6.0	(I GST-EXI GSGRE)	Converted	1/7/2005
NDA 019847			BAYER HEALTHCARE				FOR THE TREATMENT OF INHALATIONAL ANTHRAX (POST-EXPOSURE)		
Supplement 24	CIPRO	CIPROFLOXACIN	PHARMACEUTICALS	3/2/2000	8/30/2000	6.0	(I GST-EXI GSGRE)	Converted	1/7/2005
NDA 019857			BAYER HEALTHCARE				FOR THE TREATMENT OF INHALATIONAL ANTHRAX (POST-EXPOSURE)		
Supplement 27	CIPRO	CIPROFLOXACIN	PHARMACEUTICALS	3/2/2000	8/30/2000	6.0	(FOST-EXFOSURE)	Converted	1/7/2005
NDA 019858			BAYER HEALTHCARE				FOR THE TREATMENT OF INHALATIONAL ANTHRAX		
Supplement 21	CIPRO	CIPROFLOXACIN	PHARMACEUTICALS	3/2/2000	8/30/2000	6.0	(POST-EXPOSURE)	Not Converted-Application Withdrawn	6/4/2004
NDA 020780			BAYER HEALTHCARE				FOR THE TREATMENT OF INHALATIONAL ANTHRAX		
Supplement 8	CIPRO	CIPROFLOXACIN	PHARMACEUTICALS	3/2/2000	8/30/2000	6.0	(POST-EXPOSURE)	Not Converted-Application Withdrawn	6/4/2004
NDA 021174	MYLOTARG	GEMTUZUMAB OZOGAMICIN	WYETH PHARMACEUTICALS	10/29/1999	5/17/2000	6.6	FOR THE TREATMENT OF PATIENTS WITH CD33 POSITIVE ACUTE MYELOID LEUKEMIA IN FIRST RELAPSE WHO ARE 60 YEARS OF AGE OR OLDER AND WHO ARE NOT CONSIDERED CANDIDATES FOR CYTOTOXIC CHEMOTHERAPY	Not Converted-Application Withdrawn	11/28/2011
NDA 021156							TO REDUCE THE NUMBER OF ADENOMATOUS COLORECTAL POLYPS IN FAMILIAL ADENOMATOUS POLYPOSIS PATIENTS, AS AN ADJUNCT TO USUSAL		
Original 1	CELEBREX	CELECOXIB DALFOPRISTIN/QUI	GD SEARLE KING	6/25/1999	12/23/1999	6.0	CARE FOR THE TREATMENT OF VANCOMYCIN-RESISTANT	Not Converted-Indication Withdrawn	6/8/2012
NDA 050747	SYNERCID	NUPRISTIN	PHARMACEUTICALS	9/5/1997	9/21/1999	7.8†	ENTEROCOCCUS FAECIUM (VREF)	Not Converted-Application Withdrawn	11/12/2010 ²
NDA 021029	TEMODAR	TEMOZOLOMIDE	MERCK SHARP & DOHME	8/13/1998	8/11/1999	11.9	FOR THE TREATMENT OF ADULT PATIENTS WITH REFRACTORY ANAPLASTIC ASTROCYTOMA	Converted	3/15/2005
NDA 050718		DOXORUBICIN	JANSSEN				FOR THE TREATMENT OF METASTATIC CARCINOMA OF THE OVARY IN PATIENTS WITH DISEASE THAT IS REFRACTORY TO BOTH PACILTAXEL-AND PLATINUM-		
Supplement 6	DOXIL	HYDROCHLORIDE	PRODUCTS LP	12/29/1998	6/28/1999	6.0	BASED CHEMOTHERAPY REGIMENS	Converted	6/10/2008
NDA 021007	ENERASE (capsu	AMPRENAVIR	GLAXOSMITHKLINE	10/16/1998	4/15/1999	6.0	IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS, FOR THE TREATMENT OF HIV-1 INFECTION IN COMBINATION WITH OTHER ANTIRETROVIRAL	Converted	5/11/2001
NDA 021039	(oral solution)	AMPRENAVIR	GLAXOSMITHKLINE PACIRA	12/8/1998	4/15/1999	4.2	AGENTS, FOR THE TREATMENT OF HIV-1 INFECTION	Converted	5/11/2001
NDA 021041	DEPOCYT	CYTARABINE(lipos omal)	PACIRA PHARMACEUTICALS	10/5/1998	4/1/1999	5.9	FOR THE INTRATHECAL TREATMENT OF LYMPHOMATOUS MENINGITIS	Converted	4/19/2007
		DENILEUKIN					FOR THE TREATMENT OF PERSISTENT OR RECURRENT CUTANEOUS T-CELL LYMPHOMA WHOSE MALIGNANT		
BLA 103767	ONTAK	DIFTITOX	EISAI	12/9/1997	2/5/1999	13.9	CELLS EXPRESS THE CD25 COMPONENT OF THE IL-2 RECEPTOR	Converted	10/15/2008
NDA 020977	ZIAGEN (tablets)	ABACAVIR SULFATE	VIIV HEALTHCARE	6/24/1998	12/17/1998	5.8	IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS, FOR THE TREATMENT OF HIV-1 INFECTION	Converted	4/15/2004
NDA 020978	ZIAGEN (oral solution)	ABACAVIR SULFATE	VIIV HEALTHCARE	6/24/1998	12/17/1998	5.8	IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS, FOR THE TREATMENT OF HIV-1 INFECTION	Converted	4/15/2004
	1		1						

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NDA 020972	SUSTIVA	EFAVIRENZ	BRISTOL MYERS SQUIBB	6/11/1998	9/17/1998	3.2	IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS, FOR THE TREATMENT OF HIV-1 INFECTION	Converted	2/9/2000
14071 020372	00011177	LITTUINENZ	BOEHRINGER	0/11/1000	3/11/1330	0.2	PROVIDES FOR AN ORAL SUSPENSION IN COMBINATION	Convented	2/3/2000
NDA 020636			INGELHEIM				WITH OTHER ANTIRETROVIRAL AGENTS FOR THE		
Supplement 9	VIRAMUNE	NEVIRAPINE	PHARMACEUTICALS	3/16/1998	9/11/1998	5.9	TREATMENT OF HIV-1 INFECTION	Converted	3/27/2002
			BOEHRINGER				IN COMBINATION WITH OTHER ANTIRETROVIRAL		0.0
			INGELHEIM				AGENTS FOR TREATMENT OF HIV-1 INFECTION		
NDA 020933	VIRAMUNE	NEVIRAPINE	PHARMACEUTICALS	4/20/1998	9/11/1998	4.7		Converted	3/27/2002
							FOR THE TREATMENT OF MODERATELY TO SEVERELY		
							ACTIVE CROHN'S DISEASE FOR THE REDUCTION OF THE		
							SIGNS AND SYMPTOMS, IN PATIENTS WHO HAVE AN		
							INADEQUATE RESPONSE TO CONVENTIONAL		
							THERAPIES AND TREATMENT OF PATIENTS WITH		
							FISTULIZING CROHN'S DISEASE FOR THE REDUCTION IN THE NUMBER OF DRAINING ENTEROCUTANEOUS		
BLA 103772	REMICADE	INFLIXIMAB	JANSSEN BIOTECH	12/30/1997	8/24/1998	7.8	FISTULAS	Converted	4/1/2003
DENTIONIZ	KEIMIOKBE	II VI LIXIIVIX D	074400EI4 BIOTEOIT	12/00/100/	0/24/1000	7.0	FOR THE TREATMENT OF PULMONARY TUBERCULOSIS	Convented	4/1/2000
NDA 021024	PRIFTIN	RIFAPENTINE	SANOFI AVENTIS	12/22/1997	6/22/1998	6.0	FOR THE TREATMENT OF FOLIMONARY TOBERCOLOSIS	Converted	6/1/2009
							FOR USES AS ADJUNCTIVE TROPICAL ANTIMICROBIAL		
		MATERIDE	MAYL AND				AGENT TO CONTROL BACTERIAL INFECTION WHEN		
NDA 019832	SULFAMYLON	MAFENIDE ACETATE	MYLAN INSTITUTIONAL	3/31/1997	6/5/1998	14.2†	USED UNDER MOIST DRESSINGS OVER MESHED	Not Yet Converted	
14DW 013095	SULFAINITLUIN	ACEIAIE	IINOTITOTIONAL	3/31/199/	0/3/1990	14.2	AUTOGRAPFTS ON EXCISED BURN WOUNDS	Not ret Conveneu	
							FOR THE TREATMENT OF METASTATIC BREAST CANCER		
							RESISTANT TO BOTH PACILTAXEL AND AN ANTHRACYCLINE-CONTAINING CHEMOTHERAPY		
							REGIMEN OR RESISTANT TO PACILTAXEL AND FOR		
							WHOM FURTHER ANTHRACYCLINE THERAPY AY BE		
NDA 020896	XELODA	CAPECITABINE	HOFFMAN LA ROCHE	10/31/1997	4/30/1998	6.0	CONTRAINDICATED	Converted	9/7/2001
							FOR THE TREATMENT OF HIV-1 INFECTION IN		
		DELAVIRDINE					COMBINATION WITH APPROPRIATE ANTIRETROVIRAL		
NDA 020705	RESCRIPTOR	MESYLATE	VIIV HEALTHCARE	7/15/1996	4/4/1997	8.6	AGENTS WHEN THERPAY IS WARRANTED	Converted	5/16/2001
NDA 020778	VIRACEPT (oral powder)	NELFINAVIR MESYLATE	AGOURON PHARMACEUTICALS	12/26/1996	3/14/1997	2.6	FOR THE TREATMENT OF HIV INFECTION WHEN THERAPY IS WARRANTED	Converted	5/17/2000
NDA 020776	powder)	WESTLATE	FHARIVIACEUTICALS	12/20/1990	3/14/1997	2.0	FOR THE TREATMENT OF HIV INFECTION WHEN	Convented	3/17/2000
	VIRACEPT	NELFINAVIR	AGOURON				ANTIRETROVIRAL THERAPY IS WARRANTED		
NDA 020779	(tablets)	MESYLATE	PHARMACEUTICALS	12/26/1996	3/14/1997	2.6	ANTINE THOU THE TOWN THE WAR THE BOTTOM TO WAR T	Converted	5/17/2000
		MIDODRINE	SHIRE				FOR THE TREATMENT OF SYMPOTOMATIC		
NDA 019815	PROAMATINE	HYDROCHLORIDE	DEVELOPMENT	9/25/1995	9/6/1996	11.4†	ORTHOSTATIC HYPROTENSION (OH)	Not Yet Converted	
NDA 000004	OFFICATIVA	COMMEDODIN	EMD OFFICIAL	0/44/4005	0/00/4000	44.4	FOR THE TREATMENT OF AIDS WASTING AND CACHEXIA	Occupanted	0/00/0000
NDA 020604	SEROSTIM	SOMATROPIN	EMD SERONO	9/11/1995	8/23/1996	11.4		Converted	8/29/2003
			BOEHRINGER				IN COMBINATION WITH NUCLEOSIDE ANALOGUES FOR THE TREATMENT OF HIV-1 INFECTED ADULTS WHO		
			INGELHEIM				HAVE EXPREIENCED CLINICAL AND/OR IMMUNOLOGICAL		
NDA 020636	VIRAMUNE	NEVIRAPINE	PHARMACEUTICALS	2/23/1996	6/21/1996	3.9	DETERIORATION	Converted	3/27/2002
							FOR THE TREATMENT OF METASTATIC CARCINOMA OF		
		IRINOTECAN HCL					THE COLON OR RECTUM WHOSES DIEASE HAS		
NDA 020571	CAMPTOSAR	TRIHYDROTE	PFIZER	12/28/1995	6/14/1996	5.6	PROGRESSED FOLLOWING 5-FU-BASED THERAPYC	Converted	10/22/1998
							FOR THE TREATMENT OF PATIENTS WITH LOCALLY		
							ADVANCED OR METASTATIC BREAST CARCINOMA WHO		
							HAVE PROGRESSED DURING ANTHRACYCLINE-BASED		
NDA 020449	TAXOTERE	DOCETAXEL	SANOFI AVENTIS	7/27/1994	5/14/1996	21.6	THERAPY OR HAVE RELAPSED DURING ANTHRACYCLINE-BASED ADJUVANT THERAPY	Converted	6/22/1998
	., o.o. Line	200277002	2. 3.0. 17.1.2.1110	.,2.,,.004	3, 1 1, 1000	21.0	TO REDUCT CUMULATIVE RENAL TOXICITY ASSOCIATED	5511151150	3,22,1000
							WITH REPEATED ADMINSTATIONS OF CISPLATIN IN		
NDA 020221			CLINIGEN				PATIENTS WITH NON-SMALL CELL LUNG CANCER		
Supplement 2	ETHYOL	AMIFOSTINE	HEALTHCARE	2/9/1996	3/15/1996	1.2		Not Converted-Indication Withdrawn	3/28/2006 ³
NDA 000005	CDIVIVAN	INDINAVIR	MERCK SHARP &	4/04/4000	2/42/4000	4.4	FOR THE TREATMENT OF HIV-1 INFECTION IN ADULTS	Compared	0/0/4000
NDA 020685	CRIXIVAN	SULFATE	DOHME	1/31/1996	3/13/1996	1.4	WHEN THERAPY IS WARRANTED	Converted	2/6/1998
	NORVIR (oral		ABBOTT				IN COMBINATION WITH NUCLEOSIDE ANALOGS OR AS		
NDA 020659	solution)	RITONAVIR	LABORATORIES	12/21/1995	3/1/1996	2.3	MONOTHERPY FOR THE TREATMENT OF HIV INFECTION WHEN THERAPY IS WARRANTED	Converted	5/26/1999
	22.2	• . • . • . • . • . • . • . • .				1.0	IN COMBINATION WITH NUCLEOSIDE ANALOGS OR AS	2201100	2,23,1000
	NORVIR (oral		ABBOTT				MONOTHERPY FOR THE TREATMENT OF HIV INFECTION		
		RITONAVIR	LABORATORIES	12/21/1995	3/1/1996	2.3	WHEN THERAPY IS WARRANTED	Converted	5/26/1999
NDA 020680	solution)								
NDA 020680	solution)						IN COMBINATION WITH NUCLEOSIDE ANALOGS FOR THE		
NDA 020680 NDA 020628	solution) INVIRASE	SAQUINAVIR MESYLATE	HOFFMAN LA ROCHE	8/31/1995	12/6/1995	3.2	IN COMBINATION WITH NUCLEOSIDE ANALOGS FOR THE TREATMENT OF ADVANCED HIV INFECTION IN SELECTED PATIENTS	Converted	9/27/1996

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NDA 050718	DOXIL	DOXORUBICIN HYDROCHLORIDE	JANSSEN PRODUCTS LP	9/7/1994	11/17/1995	14.3	FOR THE TREATMENT OF KAPOSI'S SARCOMA IN AIDS PATIENTS WITH DISEASE THAT HAS PROGRESSED ON PRIOR COMBINATION CHEMOTHERAPY OR IN PATIENTS WHO ARE INTOLERANT TO SUCH THERAPY	Converted	6/10/2008
							IN COMBINATION WITH RETROVIR (ZIDOVUDINE) FOR THE TREATMENT OF HIV INFECTION WHEN THERAPY IS WARRANTED BASED ON CLINICAL AND/OR IMMUNOLOGICAL EVIDENCE OF DISEASE PROGRESSION		
NDA 020564	EPIVIR	LAMIVUDINE	VIIV HEALTHCARE	7/7/1995	11/17/1995	4.4		Converted	4/11/1997
NDA 020596	EPIVIR	LAMIVUDINE	VIIV HEALTHCARE	7/7/1995	11/17/1995	4.4	IIN COMBINATION WITH RETROVIR (ZIDOVUDINE) FOR THE TREATMENT OF HIV INFECTION WHEN THERAPY IS WARRANTED BASED ON CLINICAL AND/OR IMMUNOLOGICAL EVIDENCE OF DISEASE PROGRESSION	Converted	4/11/1997
							IN COMBINATION THERAPY WITH AN LHRH ANALOGUE		
NDA 020498	CASODEX	BICALUTAMIDE	ANI PHARMACEUTICALS	9/14/1994	10/4/1995	12.7	FOR THE TREATMENT OF ADVANCED PROSTATE CANCER	Converted	12/12/1997
NDA 020212	ZINECARD	DEXRAZOXANE HYDROCHLORIDE	PHARMACIA AND UPJOHN	8/5/1994	5/26/1995	9.7†	FOR THE PREVENTION OF CARDIOMYOPATHY ASSOCIATED WITH DOXORUBICIN ADMINISTRATION	Converted	10/31/2002
NDA 020412	ZERIT	STAVUDINE	BRISTOL MYERS SQUIBB	12/28/1993	6/24/1994	5.9	FOR THE TREATMENT OF ADULT PATIENTS WITH ADVANCED HIV INFECTION WHO ARE INTOLERANT OF APPROVED THERAPIES WITH PROVEN CLINICAL BENEFIT OR WHO HAVE EXPERIENCED SIGNIFICANT CLINICAL OR IMMUNOLOGICAL DETERIORATION WHILE RECEIVING THESE THERAPIES OR FOR WHOM SUCH THERAPIES ARE CONTRAINDICATED	Converted	12/21/1995
NDA 050697 Original 1	BIAXIN	CLARITHROMYCIN	ABBVIE	11/2/1992	12/23/1993	13.7	FOR THE TREATMENT OF DISSEMINATED MYCOBACTERIAL INFECTIONS DUE TO MYCOBACTERIUM AVIUM AND MYCOBACTERIUM INTRACELLULARE	Converted	5/24/2002
NDA 050698	BIAXIN	CLARITHROMYCIN	ABBVIE	11/2/1992	12/23/1993	13.7	FOR THE TREATMENT OF DISSEMINATED MYCOBACTERIAL INFECTIONS DUE TO MYCOBACTERIUM AVIUM AND MYCOBACTERIUM INTRACELLULARE	Converted	5/24/2002
BLA 103471	BETASERON	INTERFERON BETA-1B	BAYER HEALTHCARE PHARMACEUTICALS	6/18/1992	7/23/1993	13.2	USE IN AMBULATORY PATIENTS WITH RELAPSING- REMITTING MULTIPLE SCLEROSIS TO REDUCE THE FREQUENCY OF CLINICAL EXACERBATIONS	Converted	3/14/2003
NDA 020199	HIVID	ZALCITABINE	HOFFMAN LA ROCHE	10/31/1991	6/19/1992	7.6	IN COMBINATION WITH ZIDOVUDINE AS INDICATIONED FOR THE TREATMENT OF ADULT PATIENTS WITH ADVANCED HIV INFECTION (CD4 CELL COUNTS < 300 CELLS/MM3) WHO HAVE DEMONSTRATED SIGNIFICANT CLINICAL OR IMMUNOLOGIC DETERIORATION	Converted	6/26/1996

The Therapeutic Biologic Products transferred from CBER to CDER effective 1-Oct-03.

- 1. Proprietary name yet to be determined
- 2. Federal Register Date, which may be different than applicant submission or FDA action date
- 3. Date of the labeling supplement approval that removed this indication

^{†--} Total approval time was adjusted based on management decision. This is a legacy practice and is no longer exercised.