agents and antineoplastic agentsINVENTOR(S):Mittmann, Ulrich; Sachetto, Jean-PierrePATENT ASSIGNEE(S):Tillotts Pharma AG, Switz.SOURCE:PCT Int. Appl., 33 pp.
CODEN: PIXXD2DOCUMENT TYPE:PatentLANGUAGE:EnglishFAMILY ACC. NUM. COUNT:1PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.										
WO 2005123061	A1 2005122	9 WO 2005-EP6413										
		, BA, BB, BG, BR, BW,										
		, DM, DZ, EC, EE, EG,										
		, IN, IS, JP, KE, KG,										
		, MA, MD, MG, MK, MN,										
		, PL, PT, RO, RU, SC,										
		, TT, TZ, UA, UG, US,										
ZA, ZM, ZW	10, 11, 11, 11,											
	KE. LS. MW. MZ	, NA, SD, SL, SZ, TZ,	UG ZM ZW AM									
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		, CF, CG, CI, CM, GA,										
MR, NE, SN, TD, TG PRIORITY APPLN. INFO.: GB 2004-13730 A 20040618												
ED Entered STN: 30 De	c 2005	GB 2004 13730	A 20040018									
		a pharmagol accort	able salt or derivative									
			n with at least one of									
		ntineoplastic agent (
			of conditions involving									
			opical application of									
			inal mucosa. Specific									
	conditions that may be treated include chronic inflammatory disease (e.g. Chrohn's disease and ulcerative colitis) and tumor disease (e.g. bowel cancer											
			diments of the invention									
		uppressive or antine										
		contained fish oil (
		OD coating, polysorba										
REFERENCE COUNT:	,	E 10 CITED REFERENCES										
	RECORD. A	ALL CITATIONS AVAILAB	LE IN THE RE FORMAT									
L139 ANSWER 60 OF 73 HC.												
ACCESSION NUMBER:	2005:1355507 H	HCAPLUS Full-text										
DOCUMENT NUMBER:	144:74884											
TITLE:	A pharmaceutica	al compositions conta	ining									
	polyunsaturated	d fatty acids in comb	ination with									
	immunosuppress	ive agents and antine	oplastic agents									
INVENTOR(S):	Mittmann, Ulric	ch; Sachetto, Jean-Pi	erre									
PATENT ASSIGNEE(S):	Tillotts Pharma	a AG, Switz.										
SOURCE:	PCT Int. Appl.,	33 pp.										
	CODEN: PIXXD2											
DOCUMENT TYPE:	Patent											
LANGUAGE:	English											
FAMILY ACC. NUM. COUNT:	1											
PATENT INFORMATION:												

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005123060	A1	20051229	WO 2005-EP6412	20050615

	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	сн,	
		CN,	со,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
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PRIORITY APPLN. INFO.: GB 2004-13729 A 20040618																		
ED Entered STN: 30 Dec 2005																		
AB Po	lyuns	satd.	. fat	ty a	acid	("PU	FA")	or	a ph	arma	col.	acc	epta	ble	salt	or	deriv	vative
th	ereof	E (su	ich a	is EB	PA and	d/or	DHA) is	use	d in	com	bina	tion	wit	h at	lea	st or	ne of
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ac	utely	/ or	chro	onica	ally :	inad	lequa	te i	mmun	le re	spon	se b	y to	pica	l ap	plic	ation	n of –
sa	id ac	ctive	e age	ents	to a	t le	ast	а ро	ortic	n of	the	int	esti	nal	mucc	sa.	Speci	ific
co	nditi	ons	that	: may	/ be 1	trea	ted	incl	ude	chro	nic	infl	amma	tory	dis	ease	e (e.g	g.
Ch	rohn'	s di	seas	se ar	nd ul	cera	tive	col	itis) an	d tu	mor	dise	ase	(e.g	. bc	wel d	cancer
an	d pro	stat	te ca	incer	:). (One	adva	ntag	e of	pre	ferr	ed e	mbod	imen	ts c	f th	e inv	vention
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INVENTOR					Schu		-											
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SOURCE:										l., 1	ll pp	o., (Cont.	-in-	part	: of	U.S.	
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PRIORITY	APPI	LN. 1	INFO.	.:							03-4			P	20	0304	109	
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									Ľ	JS 20	04-9	7199	17	A	2 20	0410)22	
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AB Th	is in	vent	ion	prov	ides	art	icle	s an	d me	thod	s fo	r dr	ug d	eliv	ery	incl	uding	ja
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di								ns.		mpla	ry d	rugs	are	ant	i-an	giog	enesi	S

disease and/or dry eye conditions. Exemplary drugs are anti-angiogenesis compds. for the treatment of macular degeneration. Allowing passive transference of this drug from a dilute solution into the hydrogel produces the delivery system. The hydrogel, when placed in contact with the eye, delivers the drug. The delivery of the drug is sustained over an extended

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period of time, which is of particular utility in the eye, which is periodically flushed with tears. This sustained delivery accelerates the treatment process while avoiding potential damaging effects of localized delivery of high concns. of compds., e.g., from eye drops.

L139 ANSWER 62 OF 73 HC	CAPLUS	COPYRIGHT 20	006 ACS on STN										
ACCESSION NUMBER:	2005:9	83601 HCAPI	LUS Full-text										
DOCUMENT NUMBER:	143:27	2523											
TITLE:	Stable	ophthalmic	oil-in-water emulsio	ons containing									
	sodium	hyaluronate	e for alleviating dry	у еуе									
INVENTOR(S):	Yu, Zh	i-Jian; Huth	n, Stanley W.; Crawfo	ord, Lauren L.;									
	Cook,	James N.											
PATENT ASSIGNEE(S):	USA												
SOURCE: U.S. Pat. Appl. Publ., 18 pp., Contin-part of U.S.													
	Ser. No. 802,153.												
		USXXCO											
DOCUMENT TYPE:	Patent												
	Englis	h											
FAMILY ACC. NUM. COUNT:	3												
PATENT INFORMATION:													
PATENT NO.	RIND	DATE	APPLICATION NO.	האתב									
FAILNI NO.		DATE		DAIL									
US 2005196370	A1	20050908	US 2005-98827	20050404									
US 2004185068													
US 2004191284	A1	20040930											
PRIORITY APPLN. INFO.:			US 2003-392375	A2 20030318									
			US 2004-802153	A2 20040317									

ED Entered STN: 09 Sep 2005

AB Stable oil-in-water emulsions are described which contain a demulcent for the treatment of dry eye such as sodium hyaluronate. The oil-in-water emulsions are stable and have anti-microbial activity sufficient for use as contact lens disinfecting solns. Thus, an emulsion contained sodium chlorite 65 and WSCP 3 ppm, sodium hyaluronate 0.1, castor oil 1.25, ethoxylated hydrogenated castor oil 1, boric acid 0.6, sodium borate decahydrate 0.035, calcium chloride dihydrate 0.006, MgCl2.6H2O 0.006, KCl 0.14, NaCl 3.5, and water qs to 100%.

L139 ANSWER 63 OF 73 HC ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:	2004:9 141:38 Novel use Mannin Krause Biode Univer PCT In CODEN Patent Englis	902155 HCAE 84286 encochleati e-Elsmore, S livery Scien rsity of Mec nt. Appl., 1 : PIXXD2	2155 HCAPLUS <u>Full-text</u> 286 ncochleation methods, cochleates and methods , Raphael J.; Gould-Fogerite, Susan; Elsmore, Sara L.; Delmarre, David; Lu, Ruying very Sciences International, Inc., USA; ity of Medicine and Dentistry of New Jersey . Appl., 195 pp. PIXXD2						
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE					
WO 2004091578 WO 2004091578 WO 2004091578 WO 2004091578		20041028 20050127 20050331	WO 2004-US11026	20040409					
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ΕP	1624				A2			0215			004-				_	0404		
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PRIORITY	APP.	LN.	INFO	.:							003-4			-		0304		
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AB The invention generally relates to cochleate drug delivery vehicles. Disclose are novel methods for making cochleates and cochleate compns. that include introducing a cargo moiety to a liposome in the presence of a solvent. Also disclosed are cochleates and cochleate compns. that include an aggregation inhibitor, and optionally, a cargo moiety. Addnl., anhydrous cochleates that include a protonized cargo moiety, a divalent metal cation and a neg. charge lipid are disclosed. Methods of using the cochleate compns. of the invention, including methods of administration, are also disclosed.

L139 ANSWER 64 OF 73HCAPLUSCOPYRIGHT 2006 ACS on STNACCESSION NUMBER:2004:100508HCAPLUSFull-textDOCUMENT NUMBER:140:157440140:157440TITLE:Methods for treating an autoimmune disease using a soluble CTLA4 molecule in combination with a DMARD or NSAID														
INVENTOR(S):	Cohen	Cohen, Robert; Carr, Suzette; Hagerty, David; Peach, Robert J.; Becker, Jean-Claude												
PATENT ASSIGNEE(S): USA														
SOURCE: U.S. Pat. Appl. Publ., 189 pp., Contin-part of U.S. Ser. No. 898,195. CODEN: USXXCO														
DOCUMENT TYPE:	Patent													
LANGUAGE:	Englis	sh												
FAMILY ACC. NUM. COUNT: PATENT INFORMATION:	-													
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE										
US 2004022787		20040205	US 2003-419008	20030418										
US 2003083246														
PRIORITY APPLN. INFO.:			US 2000-215913P											
			US 2001-898195											
ED Entered STN: 08 Fe	eb 2004													

AB The present invention relates to compns. and methods for treating immune system diseases such as rheumatic disease, by administering to a subject

soluble CTLA4 (cytotoxic T lymphocyte antigen 4) mols. that block endogenous B7 (CD80) mols. from binding their ligands, alone, or in conjunction with other agents including disease modifying anti-rheumatic drugs (DMARDs) or nonsteroidal anti-inflammatory drugs (NSAIDs). The soluble CTLA4 mol. comprises the extracellular domain (residues 1-124) of full-length human CTLA4, which may be fused at the N-terminus with the signal peptide of oncostatin M and at the C-terminal end with an Igyl constant region. Single-site and double-site CTLA4 mutant sequences are also constructed, including L104E/A29Y-CTLA4/Ig, L104E/A29L-CTLA4/Ig, L104E/A29T-CTLA4/Ig, and L104E/A29W-CTLA4/Ig. CTLA4/Ig administered at 10 mg/kg (plus methotrexate) has superior efficacy in treatment of rheumatoid arthritis compared to placebo (plus methotrexate) based on efficacy parameters of the American Collage of Rheumatol. Core Data Set and Response Definitions (ACR). Binding kinetics to CD86 and CD80, pharmacokinetics, and pharmacodynamics of C-reactive protein, rheumatoid factor, interleukin-2 receptor, interleukin -6, and tumor necrosis factor α . are provided.

L139 ANSWER 65 OF 73 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2002:9767 HCAPLUS Full-text DOCUMENT NUMBER: 136:74627 TITLE: Drug compositions containing cyclosporin and their application as topical systems INVENTOR(S): Wohlrab, Johannes; Neubert, Reinhard; Jahn, Konstanze PATENT ASSIGNEE(S): Germany SOURCE: Ger. Offen., 14 pp. CODEN: GWXXBX DOCUMENT TYPE: Patent LANGUAGE: German FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	PATENT NO.					KIND DATE				APPLICATION NO.					DATE			
		1002 2470				A1 AA					DE 2000-10029404 CA 2001-2470230							
	WO	2003	0513	85		A1					WO 2001-EP14749							
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			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR						
	BR	2001	0171	97		А		2004	1214		BR 2	2001-3	1719	7		2	0011	214
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	ΝZ	5340	61			А		2006	0127		NZ 2	2001-	5340	61		2	0011	214
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