

No.	M/O	Label	Туре	Size	Name	Description
1	Ρ	KOD	C	7	SÚKL code	The code of the medicinal product (hereinafter referred to as "MP") allocated by SÚKL to the presentation of the MP as part of the marketing authorisation (MA) of the MP, or allocated to a non-authorised MP included in a specific therapeutic programme (hereinafter referred to as "STP") or allocated to food for special medical purposes (hereinafter referred to as "FSMP").
2	Р	NAZ	C	70	Name of the MP	The name of the MP, FSMP or STP as referred to by SÚKL guideline REG-29, version 4, in compliance with Directive 2001/83/ES.
3	Р	SILA	C	24	Strength	The strength of the MP, i.e. the contents of active substances expressed quantitatively with a view to a unit of dose, volume or weight, depending on the pharmaceutical form.
4	Р	FORMA	С	27	Pharmaceutical form	Pharmaceutical form
5	Р	BALENI	С	22	Pack	Pack size
6	Р	CESTA	С	15	Route of administration	Route of administration
7	Р	DOP	С	75	Specification of the MP	MP name supplement, which clearly defines the presentation of the MP, comprising of an integration of its pharmaceutical form, pack size, and strength. This item of the List is further specified in the items CESTA, FORMA, BALENI and SILA.
8	Р	OBAL	С	3	Packaging	The immediate packaging of the MP, i.e. such form of packaging which is in immediate contact with the MP.
9	N	DRZ	C	4	MA holder	The abbreviation for the marketing authorisation holder. A common implemental index is available for the DRZ and ZEM DRZ fields.
10	N	ZEMDRZ	С	3	Holder´s country	An abbreviation of the country of the marketing authorisation holder's registered office; for medicinal products included in specific therapeutic programmes and for foods for special medical purposes this shall mean the abbreviation of the country of the manufacturer's/importer's registered office. A common implemental index is available for the DRZ and ZEM DRZ fields.
11	N	RC	C	16	MA number	The marketing authorisation number, which identifies a group of presentations of a medicinal product for which the marketing authorisation has been issued.
12	N	SOUBDOV	C	11	Parallel import identifier	The identification number of parallel import, which is associated with the respective reference product as per the MA number; usually in the following format: PI/xxx/tyty.
13	N	T_REG	C	3	MA type	Marketing authorisation (type of marketing authorisation: national, MRP, DCP, via centralised procedure, adopted MA, parallel import).
14	Ρ	S_REG	c	2	MA status	 Status of the marketing authorisation, the basic values being as follows: B Following an implemented variation thereto, the product may be marketed for the period of 6 months and used until its expiry date, not exceeding the MA expiry date; C Revoked marketing authorisation with permitted final sale of the medicinal product; the product is to be recalled prior to the timeline specified in the decision on marketing authorisation revocation; F Specific therapeutic programme authorised by the Ministry of Health of the Czech Republic upon SÚKL's recommendation; P FSMP; R Authorised MP; Y Marketing authorisation which ceased to be valid; the product is to be recalled prior to the timeline specified in the decision.
15	Р	TCR	C	3	Price regulation type	Type of price regulation applicable values:



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						MCV Maximum ex-factory price; OP Regulation of the profit margin; the factory price is not subject to regulation under the Price Regulation of the Ministry of Health of the Czech Republic
16	N	СР	N	13,2	Producer price	The producer price (CP) of the MP/FSMP, depending on the TCR field value, this may be either the maximum ex-factory price, or the announced producer price.
17	N	LEG_CP	C	1	Producer price legal basis	 The legal basis for the determination of the producer price of a MP/FSMP/STP; it may assume the following values: G Determined <i>ex lege</i>, i.e. established by law pursuant to Section 39g, paragraph 9 of Act No 48/1997 Coll.; M Determined by the Ministry of Finance of the Czech Republic pursuant to Act No 265/1991 Coll. and Act No 526/1990 Coll. prior to 31 December 2007; N The specified price is the price determined or amended <i>ex lege</i>, for which the applicant may place the MP or FSMP on the market if their application has not been decided within timelines set forth by Act No 48/1997 Coll., as amended. This price equals the price specified in the application for maximum price determination or reimbursement. The validity of this price is limited until an enforceable decision is issued on the matter; O Announced producer price; P Temporary <i>ex lege</i> reimbursement decrease, i.e. a temporary reimbursement decrease set forth by law; R The price limit is based upon the last announced producer price, in cases where the medicinal product was re-classified by a price decision of the Ministry of Health of the Czech Republic to a maximum-price regulation, no maximum price has been determined for it to date; S Determined or amended via an administrative procedure pursuant to Act No 48/1997 Coll., as amended as of 01 January, 2008; X The decision on the maximum price has not become final as yet and is preliminarily enforceable.
18	N	ODKAZ_CP	С	20	Grounds for CP determination	If the TCR field contains the "MCV" value, this field will contain the file no. of SÚKL administrative procedure, or will be populated with the "MF" value, if the producer price has been set in compliance with existing legal regulations (price assessment of the Ministry of Finance). If the TCR field contains the "OP" value, this field will remain blank.
19	N	UHR1	N	13,2	Reimbursement	The amount of reimbursement of the medicinal product for the end consumer (JUHR1 incremented with the maximum profit margin as per the price regulation of the Ministry of Health of the Czech Republic and VAT).
20	N	JUHR1	N	13,2	Core reimbursement	The amount of reimbursement of the medicinal product determined by SÚKL as per Section 39g, paragraph 4 of Act No 48/1997 Coll. or determined as per transitory provisions of so called "technical amendment".
21	Ρ	LEG_JUHR1	C	1	Legal basis for core reimbursement	 The legal basis for the determination of the amount and conditions of core reimbursement of a MP/FSMP from health insurance; it may assume the following values: 1 The first temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; 2 The second temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; A Ex lege (statutory) reimbursement of MPs that contain an active substance listed under Section 15, paragraph 4 of Act No 48/1997 Coll., as amended in the amount of the producer price of the least



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						 economically demanding presentation; E The decision on the amount and conditions of the first temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable; F The decision on the amount and conditions of the second temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable; G Determined <i>ex lege</i>, i.e. determined by law pursuant to Section 39g, paragraph 9 of Act No 48/1997 Coll.; M Determined by Decree No 63/2007 Coll. of the Ministry of Health, the validity being governed by Act No 261/2007 Coll.; O Reduction of reimbursement as part of government-approved measure to ensure financial stability of the health insurance system as per Section 39i, paragraph 3 of Act No 48/1997 Coll.; P Temporary <i>ex lege</i> price decrease, i.e. a temporary price decrease set forth by law; Q Products whose reimbursement is directly affected by the reimbursement tender (RT), not equalling the reimbursement amount of the winning bidder of the reimbursement tender; S Established or amended via administrative procedure pursuant to Act No 48/1997 Coll., as amended as of 01 January 2008; T Permanent reimbursement tender (RT) V The MP is reimbursed within reimbursed care under Section 30 of Act No 48/1997 Coll., as amended (only for vaccines listed in Section 30 of the aforementioned Act). The amount of the reimbursement may not exceed the price of the least economically demanding presentation of the vaccine. X The decision on the amount and conditions of reimbursement has not become final to date, and it is preliminarily enforceable; Y The decision on the amount and conditions of permanent reimbursement of a highly innovative
22	N	ODKAZ_JUHR1	С	20	Grounds for core reimbursement amount determination	medicinal product has not become final to date, and it is preliminarily enforceable. Contains the file no. of SÚKL administrative procedure, or reference to statutory provision, or reference to a decree of the Ministry of Health determining the reimbursement in compliance with the aforementioned legal regulations.
23	N	LIM1	С	2	Reporting limit	The method of reporting MPs/FSMPs to the health insurance company. A common LIM status value implemental index is available for the LIM1, LIM2, and LIM3 fields.
24	N	OME1	С	40	Prescribing doctor's specialisation	The specification of prescription restriction based on the specialisation of the prescribing doctor. For a single MP/FSMP code it may assume several OME1 values. A common OME status value implemental index is available for the OME1, OME2, and OME3 fields.
25	N	IND1	С	1	Indication restriction flag	Indication restriction (P). The DETIND1 implemental index is available for the indication restriction detail (indication or clinical condition conditioning the reimbursement of the MP/FSMP). In respect of medicinal products reimbursed <i>ex lege</i> pursuant to Section 30 of Act No 48/1997 Coll., as amended, the particular provision of the Act is cited.
26	N	PUHR1	С	1	Full reimbursement flag	The full reimbursement flag may assume the following values: I the least economically demanding presentation of MPs fully reimbursed under the law; J MPs where MFC <= UHR1. Note: Final sales under the pricing regulations of the Ministry of Health of



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						 the Czech Republic are disregarded. Where final sales for a higher price in compliance with a price regulation of the Ministry of Health of the Czech Republic is applied, the applied price for the end consumer may exceed the specified amount of reimbursement; U MPs fully reimbursed under reimbursement agreements referred to under Section 39c, paragraph 2(d) of Act No 48/1997 Coll.
27	N	JUHR1_PLATDO	D	8	Temporary reimbursement expiry date	The temporary reimbursement is determined for the period of 24 or 12 months and the field is populated with the expiry date of the temporary reimbursement.
28	N	UHRPROC	N	5,2	Percentage reimbursement	The percentage of reimbursement by the insurance company set forth by a decree of the Ministry of Health of the Czech Republic.
29	N	DNC	С	1	Written arrangement on price in public interest	Identification of the maximum price (X) agreed between the health insurance company and the MA holder. If the calculated MFC is lower than the agreed maximum price announced by the health insurance company, the field will contain the "Y" value.
30	N	UHR2	N	13,2	Second reimbursement	The amount of the second reimbursement of a medicinal product/FSMP determined by SÚKL pursuant to Section 39b, paragraph 11, or Section 39d of Act No 48/1997 Coll. for the end consumer (JUHR2 incremented by the maximum profit margin as per the price regulation of the Ministry of Health of the Czech Republic and VAT).
31	N	JUHR2	N	13,2	Second core reimbursement	The amount of the second core reimbursement of a medicinal product/FSMP determined by SÚKL pursuant to Section 39b, paragraph 11, or Section 39d of Act No 48/1997 Coll.
32	N	LEG_JUHR2	C	1	Legal basis for second core reimbursement	 The legal basis for the determination of the amount and conditions of the second MP/FSMP core reimbursement from health insurance; it may assume the following values: 1 The first temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; 2 The second temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; E The decision on the amount and conditions of the first temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; F The decision on the amount and conditions of the first temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable; F The decision on the amount and conditions of the second temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable; G Determined <i>ex lege</i>, i.e. determined by law pursuant to Section 39g, paragraph 9 of Act No 48/1997 Coll.; O Reduction of reimbursement as part of government-approved measure to ensure financial stability of the health insurance system as per Section 39i, paragraph 3 of Act No 48/1997 Coll.; P Temporary <i>ex lege</i> reimbursement decrease, i.e. a temporary reimbursement decrease set forth by law; Q Products whose reimbursement is directly affected by the reimbursement tender (RT), not equalling the reimbursement amount of the winning bidder of the reimbursement tender; S Established or amended via administrative procedure pursuant to Act No 48/1997 Coll., as amended as of 01 January 2008; T Permanent reimbursement of a highly innovative medicinal product as per Section 39d, paragraph 4 of Act No 48/1997 Coll. U The winner of the reimbursement tender (RT)



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						 V The MP is reimbursed within reimbursed care under Section 30 of Act No 48/1997 Coll., as amended (only for vaccines listed in Section 30 of the aforementioned Act). The amount of the reimbursement may not exceed the price of the least economically demanding presentation of the vaccine. X The decision on the amount and conditions of reimbursement has not become final to date, and it is preliminarily enforceable; Y The decision on the amount and conditions of permanent reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable; Z Other increased reimbursement determined as per Section 39b, paragraph 11 of Act No 48/1997 Coll.
33	N	ODKAZ_JUHR2	С	20	Grounds for second reimbursement determination	Contains the file no. of SÚKL administrative procedure, or reference to statutory provision.
34	N	LIM2	С	2	Second reimbursement reporting limit	The method of reporting MPs/FSMPs to the health insurance company. A common LIM status value implemental index is available for the LIM1, LIM2, and LIM3 fields.
35	N	OME2	С	40	Prescribing doctor's specialisation	Specification of prescription restriction for the second MP/FSMP reimbursement based on the specialisation of the prescribing doctor. For a single MP/FSMP code it may assume several OME2 values. A common OME status value implemental index is available for the OME1, OME2, and OME3 fields.
36	N	IND2	С	1	Indication restriction flag	Indication restriction (P) for the second reimbursement of an MP/FSMP. The DETIND2 implemental index is available for the indication restriction detail (indication or clinical condition conditioning the second reimbursement of the MP/FSMP). In respect of medicinal products reimbursed <i>ex lege</i> pursuant to Section 30 of Act No 48/1997 Coll., as amended, the particular provision of the Act is cited.
37	N	PUHR2	С	1	Full reimbursement flag	The full reimbursement flag may assume the following values: I the least economically demanding presentation of MPs fully reimbursed under the law relevant to the ENNV2 field; J for MPs which are fully reimbursed in case MFC <= UHR2. Note: Final sales under the pricing regulations of the Ministry of Health of the Czech Republic are disregarded. Where final sales for a higher price in compliance with a price regulation of the Ministry of Health of the Czech Republic is applied, the applied price for the end consumer may exceed the specified amount of reimbursement; U MPs fully reimbursed under reimbursement agreements referred to under Section 39c, paragraph 2(d) of Act No 48/1997 Coll.
38	N	JUHR2_PLATDO	D	8	Temporary reimbursement validity specified in the second reimbursement field	The temporary reimbursement is determined for the period of 24 or 12 months and the field is populated with the expiry date of the temporary reimbursement.
39	N	UHR3	N	13,2		The amount of the third reimbursement of a medicinal product/FSMP determined by SÚKL pursuant to Section 39d of Act No 48/1997 Coll. for the end consumer (JUHR3 incremented by the maximum profit margin as per the price regulation of the Ministry of Health of the Czech Republic and VAT).
40	N	JUHR3	N	13,2	Third core reimbursement	The amount of the third core reimbursement of a medicinal product/FSMP determined by SÚKL pursuant to Section 39d of Act No 48/1997 Coll.
41		LEG_JUHR3	С	1	Legal basis for the third core reimbursement	The legal basis for the determination of the amount and conditions of the MP/FSMP core reimbursement from health insurance; it may assume the following values: 1 The first temporary reimbursement of a highly innovative medicinal product as referred to by Section



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				5122		 39d of Act No 48/1997 Coll.; 2 The second temporary reimbursement of a highly innovative medicinal product as referred to by Section 39d of Act No 48/1997 Coll.; E The decision on the amount and conditions of the first temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarilly enforceable; F The decision on the amount and conditions of the second temporary reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarilly enforceable; G Determined <i>ex lege</i>, i.e. determined by law pursuant to Section 39g, paragraph 9 of Act No 48/1997 Coll.; O Reduction of reimbursement as part of government-approved measure to ensure financial stability of the health insurance system as per Section 39i, paragraph 3 of Act No 48/1997 Coll.; P Temporary <i>ex lege</i> reimbursement is directly affected by the reimbursement tender (RT), not equalling the reimbursement amount of the winning bidder of the reimbursement tender; S Established or amended via administrative procedure pursuant to Act No 48/1997 Coll., as amended as of 01 January 2008; T Permanent reimbursement of a highly innovative medicinal product as per Section 39d, paragraph 4 of Act No 48/1997 Coll. U The winner of the reimbursement tender (RT) X The decision on the amount and conditions of reimbursement has not become final to date, and it is preliminarily enforceable; Y The decision on the amount and conditions of permanent reimbursement of a highly innovative medicinal product has not become final to date, and it is preliminarily enforceable;
42	N	ODKAZ_JUHR3	С	20	Grounds for third reimbursement determination	 Z Other increased reimbursement determined as per Section 39b, paragraph 11 of Act No 48/1997 Coll. Contains the file no. of SÚKL administrative procedure.
43	N	LIM3	С	2	Third reimbursement reporting limit	The method of reporting MPs/FSMPs to the health insurance company. A common LIM status value implemental index is available for the LIM1, LIM2, and LIM3 fields.
44	N	OME3	C	40	Prescribing doctor's specialisation	Specification of prescription restriction for the third MP/FSMP reimbursement based on the specialisation of the prescribing doctor. For a single MP/FSMP code it may assume several OME3 values. A common OME status value implemental index is available for the OME1, OME2, and OME3 fields.
45	N	IND3	C	1	Indication restriction flag	Indication restriction (P) for the third reimbursement of an MP/FSMP. The DETIND3 implemental index is available for the indication restriction detail (indication or clinical condition conditioning the third reimbursement of the MP/FSMP).
46	N	PUHR3	С	1	Full reimbursement flag	The full reimbursement flag may assume the following values J For MPs, which are fully reimbursed in case MFC <= UHR3. Note: Final sales under the pricing regulations of the Ministry of Health of the Czech Republic are disregarded. Where final sales for a higher price in compliance with a price regulation of the Ministry of Health of the Czech Republic is applied, the applied price for the end consumer may exceed the specified amount of reimbursement; U MPs fully reimbursed under reimbursement agreements referred to under Section 39c, paragraph 2(d)



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						of Act No 48/1997 Coll.
47	Ν	JUHR3_PLATDO	D	8	Third reimbursement expiry	The third reimbursement is determined for the period of 24 or 12 months. The field is populated with the
					date	expiry date of the temporary reimbursement.
48	N	RS	С	6	Reference group	The applicable reference group of the MP where the MP has been allocated a reference group by SÚKL when establishing the amount and conditions of reimbursement; it shall comprise of the applicable therapeutic group (TS), separator (/), sequence of the subgroup of products that are similar or that can cause confusion in the reference group (RS_P); the RS stipulated by a decree of the Ministry of Health of the Czech Republic under authority referred to under Section 39c, paragraph 1 of Act No 48/1997 Coll.
49	N	TS	N	3	Therapeutic group	The applicable therapeutic group of the MP/FSMP if it has been allocated a therapeutic group by SÚKL when establishing the amount and conditions of reimbursement; the TS stipulated by a decree of the Ministry of Health of the Czech Republic under authority referred to under Section 39c, paragraph 1 of Act No 48/1997 Coll.
50	N	TS_P	N	2	TS subgroup	The applicable subgroup of products that are similar or that can cause confusion within the TS of the MP/FSMP where an RS has been allocated by SÚKL when establishing the amount and conditions of reimbursement; the TS_P stipulated by a decree of the Ministry of Health of the Czech Republic under authority referred to under Section 39c, paragraph 1 of Act No 48/1997 Coll.
51	N	ATC	с	7	Full ATC	Anatomical therapeutic chemical group. An ATC implemental index is available for the ATC field.
52	N	V_PLATOD	D	8	MA effective date	The effective date of the marketing authorisation.
53	N	V_PLATDO	D	8	MA expiry date	The expiry date of the marketing authorisation, unless unlimited validity has been granted pursuant to Section 34 of the Act on Pharmaceuticals.
54	Ν	NEOMEZ	С	1	Unlimited MA validity	Field to be completed (X) where unlimited validity of the marketing authorisation applies.
55	N	HL_UV_OD	D	8	Placement on the market	Date of initial placement of supplies of the medicinal product on the market or reinstitution thereof, to be reported by the MA holder in compliance with Section 33 of the Act on Pharmaceuticals.
56	N	HL_UK_DO	D	8	Supply termination	Date of termination or discontinuation of supplies of the medicinal product onto the market, to be reported by the MA holder in compliance with Section 33 of the Act on Pharmaceuticals.
57	N	DDDM	С	4	Amount of active substance in DDD	Defined daily dose – the amount of active substance – information as per WHO.
58	N	DDDJ	С	2	Unit of active substance amount in DDD	Defined daily dose unit – information as per WHO.
59	Ν	DDDBAL	Ν	11,4	DDD count in MP pack	The number of defined daily doses in a pack where DDD has been established by WHO.
60	N	ODTD1	N	13,4	Usual daily therapeutic dose for reimbursement	The usual daily therapeutic dose for reimbursement.
61	N	ODTDJ1	С	5	Unit of active substance amount in ODTD1	The usual daily therapeutic dose for reimbursement unit for ODTD1.
62	N	ODTDBAL1	N	11,4	Number of ODTD1 in a MP pack	The number of usual therapeutic doses in a pack for ODTD1.
63	N	ODTD2	N	13,4	Usual daily	The usual daily therapeutic dose for second reimbursement.



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					therapeutic dose for second	
-					reimbursement	
64	Ν	ODTDJ2	С	5		The usual daily therapeutic dose for second reimbursement unit for ODTD2.
					substance amount in	
65		00700410			ODTD2	
65	N	ODTDBAL2	Ν	11,4	Number of ODTD2 in a MP	The number of usual therapeutic doses in a pack for ODTD2.
66	N	ODTD3	N	12.4	pack Usual daily	The usual daily therapeutic dose for third reimbursement.
00	IN	00103	IN	13,4	therapeutic dose for third	
					reimbursement	
67	N	ODTDJ3	с	5		The usual daily therapeutic dose for third reimbursement unit for ODTD3.
07		001033	C	5	substance amount in	The usual daily therapeatic abserior time reinbarsement and for obrids.
					ODTD3	
68	N	ODTDBAL3	N	11,4	Number of ODTD3 in a MP	The number of usual therapeutic doses in a pack for ODTD3.
				,	pack	
69	N	EKV1	N	13,4	Reimbursement for ODTD1	Basic reimbursement of the active substance or reference group or MP, resp., in case of a temporary
						reimbursement for ODTD1, if it has been determined by SÚKL pursuant to Section 39c, paragraph 1 and
						paragraph 2, or Section 39d, resp., of Act No 48/1997 Coll.
70	Ν	ODKAZ_EKV1	С	20	Grounds for EKV1	Contains the file no. of SÚKL administrative procedure.
					determination	
71	N	EKV2	Ν	13,4	Reimbursement for ODTD2	The second reimbursement of the active substance or pseudo-reference group or MP, resp., in case of a
						temporary reimbursement for ODTD2, if it has been determined by SÚKL pursuant to Section 39b, paragraph
						11, or Section 39d, resp., of Act No 48/1997 Coll.
72	Ν	ODKAZ_EKV2	С	20	Grounds for EKV2	Contains the file no. of SÚKL administrative procedure.
		_			determination	
73	Ν	EKV3	Ν	13,4	Reimbursement for ODTD3	Third reimbursement of a MP for ODTD3, if it has been determined by SÚKL pursuant to Section 39d of Act
74				20		No 48/1997 Coll.
74	N	ODKAZ_EKV3	С	20	Grounds for EKV3	Contains the file no. of SÚKL administrative procedure.
75	N	DAT CP	D	0	determination CP validity	The effective date of a producer price change.
75 76	P	DAT_CP DAT_UHR	D		UHR validity	The date of change of the determination of the amount and conditions of reimbursement.
70	P	ZAP1	N		Eligible extra	An eligible extra payment for UHR1 under a communication of the Ministry of Health of the Czech Republic,
//	P	ZAPI	IN	13,2	payment	stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997 Coll.
78	Р	NEZAP1	С	1	Limit eligibility	A flag of the category of the pharmaceutical (eligibility for the limit) under a communication of the Ministry
70	г	NLZAF I	C	T	symbol	of Health of the Czech Republic, stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997
					Symbol	Coll. The NEZAP implemental index is available for the NEZAP1 field.
79	N	ZAKL ZAP1	С	1	Eligible extra	Contains the "X" value only for MPs that form the basis for the calculation of the eligible extra payment
				1	payment calculation	under Section 16b, paragraph 1 of Act No 48/1997 Coll.
					base	
80	N	ZAP2	N	13,2	Eligible extra payment for	An eligible extra payment for UHR2 under a communication of the Ministry of Health of the Czech Republic,



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					second reimbursement	stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997 Coll.
81	N	NEZAP2	С	1	Limit eligibility symbol	A flag of the category of the pharmaceutical (eligibility for the limit) under a communication of the Ministry of Health of the Czech Republic, stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997 Coll. The NEZAP implemental index is available for the NEZAP2 field.
82	N	ZAP3	N	13,2	Eligible extra payment for third reimbursement	An eligible extra payment for UHR3 under a communication of the Ministry of Health of the Czech Republic, stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997 Coll.
83	N	NEZAP3	С	1	Limit eligibility symbol	A flag of the category of the pharmaceutical (eligibility for the limit) under a communication of the Ministry of Health of the Czech Republic, stipulated in compliance with Section 16b, paragraph 1 of Act No 48/1997 Coll. The NEZAP implemental index is available for the NEZAP3 field.
84	N	ZPVYD	C	1	Method of dispensing	 Classification of the medicinal product for dispensing: F The medicinal product may be dispensed without medical prescription; L The medicinal product may be dispensed on the basis of a medical prescription prescribed by a doctor of a specialised competence, only for providers of healthcare services providing healthcare services in inpatient care setting; O The medicinal product may be dispensed without medical prescription, but a restriction on the dispensing has been set (Section 39, paragraph 5 of the Act on Pharmaceuticals); P The medicinal product which may be dispensed to a single patient within a predefined period of time has been set (Section 39, paragraph 5 of the Act on Pharmaceuticals); R The medicinal product may be dispensed on medical prescription only; V Selected medicinal product. In case of foods for special medical purposes, the method of dispensing is not specified, as decision-making on the method of FSMP dispensing is not within the powers of SÚKL.
85	N	ENNV1	N	13,2	Economically least demanding vaccine presentation	The amount of reimbursement for the economically least demanding presentation of a vaccine reimbursed pursuant to Section 15, paragraph 4, or Section 30 of Act No 48/1997 Coll., as amended.
86	N	MFC	N	13,2	Final price	Final price – the price for the end consumer (producer price plus the maximum profit margin under the price regulation of the Ministry of Health and VAT). If the insurance company and the MA holder concluded an agreement on the maximum agreed price for the product or assumed a valid obligation not to exceed the price assumed in a pricing tender under Section 39 of Act No 48/1997 Coll., as amended prior to the technical amendment, the agreed price, lower than the MFC, will be specified in this field. If the calculated MFC is lower than the agreed maximum price announced by the insurance company or the price assumed in the pricing tender, the calculated MFC value will be specified in this field.
87	N	POCDAV	N	13,0	Number of vaccine doses	The number of vaccine doses in a pack.
88	Ν	RP1	С	1		Reserve field 1
89	Ν	RP6	С	15		Reserve field 6
90	Ν	RP7	С	10		Reserve field 7
91	Ν	DPH	Ν	13,0	Value-added tax rate	Contains a numerically expressed value-added tax rate.



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92	N	ENNV2	N	13,2	Economically least demanding vaccine presentation	The amount of reimbursement for the economically least demanding presentation of a vaccine reimbursed pursuant to Section 15, paragraph 4 or Section 30 of Act No 48/1997 Coll., as amended.
93	Ν	RP10	С	1		Reserve field 10
94	Ν	RP11	С	20		Reserve field 11
95	Ν	RP12	С	2		Reserve field 12
96	Р	NAZ_REG	С	70	Name of the MP	Authorised name of the MP/FSMP/STP
97	Ν	RP14	С	1		Reserve field 14
98	Ν	RP15	С	1		Reserve field 15
99	Ν	RP16	D	8		Reserve field 16
100	Р	POC_UHR	N	13,4	Total number of reimbursements	Total number of reimbursements established MP/FSMP
101	N	RP18	С	5		Reserve field 18
102	N	RP19	N	11,4		Reserve field 19
103	Ν	RP20	N	13,4		Reserve field 20
104	Ν	RP21	С	20		Reserve field 21
105	Ν	RP22	N	13,2		Reserve field 22
106	Ν	RP23	С	1		Reserve field 23
107	Ν	RP24	N	13,2		Reserve field 24
108	Ν	RP25	N	13,2		Reserve field 25
109	Ν	RP26	С	1		Reserve field 26
110	Ν	RP27	С	20		Reserve field 27
111	Ν	RP28	С	2		Reserve field 28
112	Ν	RP29	С	40		Reserve field 29
113	Ν	RP30	С	1		Reserve field 30
114	Ν	RP31	С	1		Reserve field 31
115	Ν	RP32	D	8		Reserve field 32
116	Ν	RP33	N	13,4		Reserve field 33
117	Ν	RP34	С	5		Reserve field 34
118	Ν	RP35	Ν	11,4		Reserve field 35
119	Ν	RP36	Ν	13,4		Reserve field 36
120	Ν	RP37	С	20		Reserve field 37
121	Ν	RP38	Ν	13,2		Reserve field 38
122	Ν	RP39	С	1		Reserve field 39

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Field separator "I"

The "M/O" column identifies mandatory and optional fields in the List The "Type" column identifies the format of the fields as follows:

"C" character attribute "N" numeric attribute



"D" date in the "ddmmyyyy" format The "Size" column identifies the scope of the fields. The format of numeric fields is identified as "x,y" ("x" positions, incl. the decimal point, of which "y" are decimals)