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¹ Initial Digestive Potential of Alimentary System in Newborns

 Penzhoyan Grigorii Artemovich¹, Model Galina Yurievna² and Korotko Gennadii Feodosyevich³
 ¹ Kuban State Medical University
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7 Abstract

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⁸ Fatally organized alimentary system hydrolase activities in a newborn make up the initial

⁹ digestive polyenzyme potential, which provides breast milk lacto trophy if combined with

 $_{10}$ $\,$ hydrolases. Initial digestive potential in a newborn is characterized by the results of the

activity and content of lipase, ?amylase, pepsinogens (I, II), alkaline phosphatase,

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¹³ newborn at the end of the delivery.Systems of different hydrolases during antenatal life are

¹⁴ asynchronous.According to the results of the hydrolase estimation in the blood serum of the

¹⁵ mother and the newborn, the digestive potential of the latter turns out to be much less than

16 that of the mother's. It is the proof of the incomplete maturity of the digestive potential in

¹⁷ the newborn. In the case of immature gestation, the concentration of hydrolases and

¹⁸ zymogens (except lipase) in the examined bio liquids was reduced. Hydrolases of gastric ¹⁹ contents are most informative towards the digestive potential and less informative towards

²⁰ amniotic fluids and umbilical cord blood serum.

22 Index terms— newborn, hydrolases, amniotic fluids, gastric content, blood serum, digestive potential.

²³ 1 Initial Digestive Potential of Alimentary System in Newborns

24 Penzhoyan G.A. ? , Model G.Y. ? & Korotko G.F. ?

Abstract-Fatally organized alimentary system hydrolase activities in a newborn make up the initial digestive polyenzyme potential, which provides breast milk lacto trophy if combined with hydrolases. Initial digestive potential in a newborn is characterized by the results of the activity and content of lipase, ?-amylase, pepsinogens (I, II), alkaline phosphatase, ? 1 -antitrypsin in umbilical cord blood serum, amniotic fluid, aspirate gastric content in a newborn at the end of the delivery. Systems of different hydrolases during antenatal life are asynchronous.

According to the results of the hydrolase estimation in the blood serum of the mother and the newborn, the digestive potential of the latter turns out to be much less than that of the mother's. It is the proof of the incomplete maturity of the digestive potential in the newborn. In the case of immature gestation, the concentration of hydrolases and zymogens (except lipase) in the examined bio liquids was reduced. Hydrolases of gastric contents are most informative towards the digestive potential and less informative towards amniotic fluids and umbilical cord blood serum.

Identification of the enzymes in three mentioned bio liquids of the newborn is advisable for the reasonable estimation of the digestive potential and precise prognosis of lacto trophy.

40 Quantitative indicator of the potential was measured by analyzing lipase, ?-amylase, alkaline phosphatase,

⁴¹ pepsinogens (I, II) in umbilical cord blood serum, amniotic fluid, aspirate gastric content in a newborn at the end ⁴² of the delivery, in venous blood of 36 new mothers with full-term pregnancy, in 40 new mothers with incomplete ⁴³ restation as well as in their newborns.

43 gestation as well as in their newborns.

<sup>Resume: Initial enzyme digestive potential is presented by fatally organized alimentary system hydrolase
activities in a newborn. Both lacto trophy efficacy and breast milk hydrolases depend on it.</sup>

During antenatal life systems of different hydrolases were formed asynchronously. In the case of immature gestation, the concentration of hydrolases and zymogens in the examined bio liquids was reduced except lipase, which was active. Hydrolases of gastric contents are most informative towards the digestive potential and its low variability, while hydrolases in the newborn umbilical cord blood serum and in the amniotic fluids are less informative. Poly-enzyme analysis of three bio-liquids of a newborn proved to be reasonable for the conclusion concerning the morph functional maturity of the digestive system of a newborn.

50 Initial digestive potential makes it possible to expect lacto trophy efficacy and individualize the program for 51 both breast and mixed feeding.

⁵² 2 I. Introduction

reastfeeding proved to be the "gold standard" for a newborn thanks to its unique nutritional, immune, regulatory, 53 electrolyte, vitamin characteristics, as well as to the microbiota of the breast milk and its numerous substitutes. 54 Milk nutrients are ingested by a newborn, but first they need to be hydrolyzed in the digestive system, this process 55 is performed by hydrolyzing enzymes of the digestive glands and small intestine of a newborn according to the 56 self-digestion pattern, and by first milk and mature milk enzymes according to autolytic digestion. Autolysis 57 of lipids and proteins (casein) of first milk and milk is induced and realized in the gastric and small intestine 58 cavities by the hydrolases of the newborn's digestive glands that developed during his antenatal period. Their 59 hydrolases make up the initial digestive potential of the alimentary tract of the newborn. The digestive potential 60 has not yet been investigated in perinatology, neonatology, or pediatrics either in terms of theory or in applied 61 medicine. The notion has recently been put forward by the authors. But it should be taken into consideration 62 that the reducing of this morphofunctional potential may threaten the development of a newborn. 63

⁶⁴ **3** II. Materials and Methods

Among seventy-six examined new mothers 36 had full-term (37-41 weeks) and 40 premature (27-36 weeks) 65 pregnancies. Forty-seven children were born during vaginal birth and 29 by cesarean section. The investigation 66 began after the written consent was signed by the parents under the current Federal "Law on Health Protection 67 of Citizens" and the decision of the Ethics Committee. In newborns, anthropometric data, Apgar score and 68 some anthropometric parameters and obstetric history were assessed under the Order of the Ministry of Health 69 of the Russian Federation "On approval of the Order of Medical Care in the Profile «Neonatology»". The 70 above-mentioned parameters were significantly lower in premature newborns than in mature newborn infants 71 (see Table 1). New mothers' amniotic fluids were obtained in sterile syringes and then centrifuged (10 min., 3000 72 revolutions). The newborns' blood was obtained from their umbilical cords; the mothers' blood was obtained 73 from the ulnar vein. 74 75 In newborns fasting gastric content was aspirated, then it was homogenized and centrifuged (10 min., 3000 76 revolutions). In amniotic fluid and gastric aspirate supernatants, umbilical cord blood serum of the newborn and mother's blood serum lipase, ?-amylase, alkaline phosphatase were determined by colorimetric methods with 77

and model is blocd solution inpuss, it any fact, and fact, and incorporation with a standard reagent kits for in vitro diagnostics (Roche) on a modular platform for biochemical and immunochemical
analysis Cobas-8000 (module C 702). ?-1-antitrypsin (reagent F1-Antitrypsin) was determined on a biochemical
analyzer Architect C 8000 (Abbott) by the turbidimetry method. Pepsinogens I and II were determined by
chemiluminescent immunoassay analysis on microparticles by Abbott reagents using immunological analyzer
Architecr plus: 2000.

Statistical data processing was implemented within the Statistica 6 package by nonparametric statistics
 methods since the above-mentioned parameters had a large spread, and their empirical values did not correspond
 to the standard distribution law. Correlation analysis of enzyme parameters was carried out.

⁸⁶ 4 III. Results and Discussion

The aim of the research is the quantitative characteristics of the initial digestive potential of the alimentary tract in newborns and methods for its determination that includes the determination of digestive glands hydrolyzes in the newborn's blood serum, gastric aspirate, and amniotic fluids.

The amount of digestive glands hydrolases in human blood serum depends on the number and activity of 90 glands producers granulocytes of the correlative enzymes [9]. In the blood serum of new mothers, the amount of 91 hydrolases is higher (Table 2) than that in the blood serum of the newborns (Table 3 It proves the incomplete 92 development of the digestive glands enzymatic potential. Hydrolases differ in the initial level of their content in 93 94 the blood serum that indicates that morphofunctional maturation of the enzyme systems of the fetus and the 95 newborn is asynchronous. Producers of pepsinogen -the stomach glands (especially pepsinogen I) and producers 96 of ?amylase -salivary and pancreas glands are most retarded. The antitrypsin activity of umbilical cord blood serum in the newborns was 4.5 times as high as that of the mother's. The new mothers that gave birth to 97 both premature and mature newborns did not differ in the content of blood serum enzymes, except for alkaline 98 phosphatase. 99

The concentration of amylase and pepsinogen (I, II) in the umbilical cord blood serum of the premature newborns was lower than that in the umbilical cord blood serum of mature newborns (Table 3). Reduced amyl lytic activity of the glands secrets in premature newborns can cause maldigestion in the case of mixed and artificial

feeding of infants as most infant formula milk contains ?-amylase-hydrolyzed polysaccharides. It is not contained 103 in breast milk. Incomplete gestation reduces premature peptic potential of the fund-antroduodenal producers 104 of pepsinogens. It may affect the hydrolysis and protein metabolism in premature newborns, the formation of 105 regulatory peptides (mainly breast milk casein) [5][6][7][8][9][10][11][12][13], and the process of proteolysis in the 106 lacto trophy. This statement results from the postulate of the interaction of breast milk proteases and digestive 107 glands excretions in the gastrointestinal tract [6,7,10], that have recently been confirmed by peptidomics and 108 mass spectral chromatography. According to this fact the excretions proteinases (gastric aspirate) increase the 109 hydrolytic effect of breast milk proteinases of lactating women by 1.5 -2.5 times [10]. The milk proteinases (like 110 other hydrolases) have specific self-regulating dynamics during lactation [1,14]. 111

Premature birth did not affect the lipase content in the umbilical blood serum in the newborns. It speaks for the formation of the low initial level of lipase production by the digestive glands of the fetus during earlier gestational periods than other considered enzymes.

The reduced content of three hydrolases (?amylase, pepsinogen I and II) in the umbilical cord blood serum of 115 the newborn proved immature initial digestive potential in preterm pregnancy. No significant data concerning 116 other hydrolases were found. It is explained by the fact that enzymatic homeostasis in the blood is provided 117 not only by transporting the corresponding enzymes and zymogens using increment and resorption but removing 118 119 the same enzymes from the bloodstream by different mechanisms. It has been the subject of quite a number 120 of experimental and clinical research (Review: [5,16]). Therefore, the relatively constant content of hydrolases at one or another level is the result of the balance of the given complex multidirectional regulated processes. 121 In newborns we observed three low hydrolases content in the blood and their severe vibrations, that prove low 122 enzyme potential, its variability, and, consequently, limited diagnostic information value. 123

¹²⁴ 5 The Digestive Glands Hydrolases in Amniotic Fluids

The volume and composition of amniotic fluids have been studied under normal and pathological conditions by 125 lots of researchers at different times. The presence of enzymes in the amniotic fluids, including digestive gland 126 hydrolases, has been established. However, the informational hydrolases criteria concerning the enzyme potential 127 of the glands have not been studied in full, especially the mechanisms of origin of this group of enzymes in the 128 amniotic fluids [15]. In different periods of gestation, hydrolases in the amniotic fluid are of different origin, but 129 at the end of the gestation they come mainly from the digestive glands of the fetus. We cannot deny participation 130 of hydrolases of amniotic fluid and placenta in the genesis [17], as well as the transport of hydrolases from the 131 blood of a pregnant woman [15,17]. They seem to be additional sources of enzymes in the amniotic fluids. These 132 problems have recently been under our consideration [14]. 133

In the amniotic fluids of new mothers with fullterm gestation, the composition of hydrolases (see Table 4) 134 differs from that in the blood serum of newborns (Table 3) and their new mothers (Table 2). The concentration 135 of ?-amylase, ?-1-antitrypsin, pepsinogen I and especially pepsinogen II in the amniotic fluids is much higher than 136 that in the blood serum of new mothers. What concerns alkaline phosphatase the differences are insignificant. 137 The lipase composition in the amniotic fluid is five times as low as in the umbilical cord blood serum and even 138 15 times as low as the average blood serum index of the new mother. We cannot but mention the moderate 139 statistically significant correlation between the hydrolase content in amniotic fluids and blood serum of umbilical 140 cord: for ?-amylase r=0,63; for pepsinogen I r=0,68; for pepsinogen II r=0,50; for alkaline phosphatase r=0,52. 141 Hence, the hydrolases of amniotic fluids are informative concerning the individual morphofunctional immaturity 142 of digestive glands in both fetuses and newborns. 143

In incomplete pregnancy, the amniotic fluids contain all types of hydrolases except lipase in a less concentration than in full-term pregnancy. The decreased content of hydrolases is statistically highly significant (p<0, 01).

High concentration of pepsinogen II in amniotic fluids that differ greatly from pepsinogen I prove the differences 146 in development mechanisms of hydrolases of digestive glands in the systemic bloodstream of newborns, their 147 mothers, and amniotic fluids. This phenomenon can be explained by the early development of enteric enzyme 148 producers in the fetus [3,[18][19][20]. Pepsinogen II is mostly synthesized by pyloric and duodenal glands. That 149 is why the concentration of this isoproenzyme in amniotic fluids in incomplete pregnancy as well. So, the 150 regurgitated stomach content is transported to the amniotic fluids as the result of the duodenal, gastric and 151 oral reflux which is common for both fetuses and newborns, while in their stomach content we found out higher 152 concentration of pepsinogen II in comparison with pepsinogen I. Evidences of these differences are given in Table 153 154 5.

The high hydrolytic activity of amniotic fluids, the high volume of their transfer into the digestive tract of the fetus by swallowing, breathing and inhaling makes it possible to conclude that hydrolases of amniotic fluids take part in the hydrolysis of nutrients of the gastrointestinal tract which provides the amniotic trophism with its specific autolytic and self-digestion. It is necessary for the nutrition of the digestive tract mucous coat structures.

¹⁵⁹ 6 Hydrolases of Aspirated Stomach Content of Newborns

Fasting stomach content of a newborn is a mixture of gastric glands duodenal contents (pancreas secretion, duodenal secretions, and bile secretions), swallowed oral liquid (secretions of salivary glands and crevicular fluids) and amniotic fluids. Due to the absence of recurring activities of the digestive system in neonates [1],

the volume and composition of the aspirated stomach content are relatively stable and demonstrate the total 163 secretory activity of the above mentioned digestive glands, including their enzyme production. Judging by the 164 data given in Table 5, polysecretion aspirated from the stomach possessed the high concentration of ?-amylase, 165 lipase and pepsinogens, especially pepsinogen II; all 76 samples of stomach content demonstrated the higher 166 level of pepsinogen II than pepsinogen I. The same result was received after the analysis of amniotic fluids. The 167 level of similar hydrolases both in the gastric aspirate and amniotic fluids had moderate statistically significant 168 correlation coefficients: for anylase r=0.57; for pepsinogen II r=0.60. We registered a strong correlation among 169 five enzymes of amniotic fluids and gastric aspirate: the index of canonical correlation was R pcc =0.82 (that 170 characterizes the stage and interaction force between two variable lists). The results of these findings prove the 171 above-formulated discovery of one more physiological mechanism, namely the development of digestive glands 172 hydrolases of high concentration in amniotic fluids: duodenogastrooral regurgitation (reflux) into the amnion. 173

High enzyme activity of gastrointestinal contents provided by the fetal enzymes of both digestive glands and enterocytes performs the cavitary, parietal, and intracellular digestion of the fetus, including its amniotic trophism. In neonatal and subsequent stages of the child's development the hydrolases of his digestive tract that made up his initial digestive potential provide (together with the breast milk hydrolases) the lacto trophy with its peculiar proper and autolytic types (including the induced subtype) of digestion.

179 Saliva proteases increase the activity of casein by pepsins and trypsin in vitro. Similar of proteinases in 180 lacto trophy takes place in the stomach and small intestine under appropriate conditions (pH of the medium) 181 [1]. In several recent works devoted to enzyme peptidomics, the summing up of the proteolysis produced by 182 secretory proteases in the baby's stomach and similar proteases of mother's milk incubated in the stomach (2 h) by nano-chromatographic identification of peptides formed mainly during hydrolysis of ? casein was established. 183 At the same time, the effects of plasmin did not change, or they reduced by 1.3 times. Cathepsin D actions 184 increased by 2.3 times, of pepsin by times, of elastase by 1.6 times, of chymotrypsin by 2.5 times, and those 185 of prolineendopeptidases by 1.5 times. Hence, milk autoproteolysis was increased twice as much by secretory 186 proteases in the stomach of the infant by the proteases [16]. The authors verified the relevance of intragastric 187 proteolysis in the formation of regulatory peptides, most of which have acknowledged effects. 188

Pediatricians take an interest in the lipolytic activity of milk and its lipids, which play energetic, plastic, 189 nutritional and protective role in the lacto trophy of the child. The lipolysis technology is multistage: it is 190 performed by lipases of saliva and gastric secretion in the stomach cavity, then by lipases of milk and pancreatic 191 secretion in the small intestine with the participation of bile salts inducers (promoting milk lipase) and colipase 192 (promoting the effect of pancreatic lipase) [15]. Triglycerides are released from milk fat globules in the stomach 193 by hydrophobic lipases of saliva and gastric secretion, that act as inducers of lingual and gastric lipases of the 194 infant as well. The material of the globules membranes is recognized as a valuable product for the infant and has 195 recently been added to milk mixtures. By the way, during the period of lactation, the lipolytic activity of milk 196 is reducing more slowly than the content of other hydrolases in milk [15]. 197

In human breast milk, there is no substrate for ?-amylase, but its activity is high in the gastric aspirate. It 198 is significant for the polysaccharide hydrolysis in complementary foods in the case of mixed and artificial feeding 199 of infants. Hydrolysis of the principal carbohydrate of lactose milk is carried out by milk lactases and the small 200 intestinal mucosa. Lots of researchers have lately focused their attention on these enzymes. Lactase is one of the 201 disaccharides of enteric membrane digestion it was not included in the secretory potential and was not found in 202 the gastric aspirate. The results shown in Table 5 indicate a significant decrease in hydrolases content (except 203 lipase and antitrypsin) in the gastric aspirate of premature newborns if compared to full-term newborns. These 204 data are extremely informative about the secretory digestive potential of newborns. 205

206 7 IV. Summary

The technology of lacto trophy makes it possible to conclude that the secretory hydrolases of the digestive glands, which form the digestive potential of the newborn, are of fundamental importance for its implementation. In this regard, its quantitative characteristic should be taken into account not only in incomplete gestation periods, but also in normal ones.

It is all the more important because hydrolase levels proved to be higher than the average in gastric aspirate, 211 amniotic fluid, and umbilical cord blood serum in the group of premature infants, who had mainly a reduced 212 213 digestive potential, while in infants of the group with standard gestational age enzymatic indicators of three 214 bio liquids were reduced in comparison with average values. This phenomenon took place at the gestational 215 borderline. Such results make it possible to acknowledge the digestive potential of newborns during childbirth 216 the diagnostic test in the trophological prognosis of the development of newborns. Due to its digital variability and quantitative insufficiency, the material obtained does not allow determining the reference enzyme parameters 217 of the standard initial digestive potential. That is why further fact-finding inquiry is necessary. At the current 218 state of knowledge only a sharp decrease in the quantitative initial digestive potential of hydrolases in amniotic 219 fluids and in umbilical cord blood serum can serve a reliable prognostic sign of trophological dysfunction in a 220 newborn. 221

222 8 V. Conclusions

1. Hydrolases of the secrets of the digestive glands and small intestine of newborns make up the prenatally formed initial digestive potential of their digestive system. 2. The digestive potential characterized by the enzymes of the cord blood serum of the infant is significantly lower than that of the mother's venous blood serum and proves the incompleteness of the digestive potential in the antenatal period.

227 9 Low enzymatic activity of this potential requires

228 proper and autolytic digestion of breast milk hydrolases to participate in lacto trophy. 4. Morphofunctional

²²⁹ maturation of producers of different digestive system hydrolases of the fetus and the child are asynchronous: the digestive system of the small intestine matures earlier than the others, next come to the lipase producing ¹

1	1	L		
		L		

Variables	Average	Median	Minimu	m Maximu	mLower	Upper	Shift
	0	value			quartile	quartile	direction,
					-	-	statistical
							significance
Mother's age	30,75	31,5	19,0	44,0	27,5	35,0	p < 0.5
(years)	$27,\!67$	28,0	14,0	41,0	24,5	$_{30,5}$	
Gestional age	32,00	33,0	27,0	35,0	30,0	34,0	$\hat{a}??$ " p < 0,001
(weeks)	38,03	38,0	38,0	39,0	38,0	38,0	
Mass	3546,4	3595,0	2460,0	4800,0	3170,0	3835,0	â??" p <
(g)	1765,2	1715,0	670,0	3130,0	1335,0	2150,0	0,001
Height	$53,\!47$	54,0	46,0	59,0	52,0	55,5	â??" p <
(cm)	$41,\!27$	42,0	33,0	48,0	37,0	46,0	0,001
Head circumfer-	$28,\!58$	30,0	17,0	35,0	26,0	31,0	â??" p < 0,001
ence (cm)	$34,\!14$	34,0	31,0	37,0	33,0	$35,\!5$	
Breast circum-	$26,\!28$	27,0	16,0	33,0	24,0	29,0	$\hat{a}??$ " p < 0,001
ference (cm)	$33,\!31$	33,0	27,0	37,0	32,0	35,0	
Apgar 1	7,9	8,0	7,0	8,0	8,0	8,0	â??" p <
(scores)	$5,\!5$	6,0	1,0	7,0	$5,\!0$	$6,\!0$	0,001
Apgar 5							
(scores)	6,0						

Figure 1: Table 1 :

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[Note: Note: ALP -alkaline phosphatase. (numerator -mature newborns, denominator -premature newborns)]

Figure 2: Table 2 :

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Enzymes	Average	Median Value	Minimum	Maximum	Lower Quartile	Upper Quartile	Shift direction Statistical significanc	e
Lipase (U/l)	$10,\!49$	10,00	$5,\!40$	18,80	$8,\!308,\!30$	12,60	p > 0,10	
	10,72	$10,\!12$	5,70	$21,\!30$		$12,\!10$		
Amylase	4,72	4,00 8,50	$0,\!00$	16,00	$2,00\ 5,00$	6,00 9,00	$\hat{a}??"p < 0$,01
(U/l)	9,00		1,00	52,00				
ALP (U/l)	166,5	171,0	$11,\!0$	274,0	134,0	200,5	p > 0, 10	
	157,5	157,5	97,0	243,0	119,0	181,5		
Pepsinogen I	4,82	$4,20\ 8,45$	$1,\!10$	12,20	$2,60\ 6,75$	$6,\!10$	$\hat{a}??$ "p	<
(ng/ml)	10,92		$3,\!40$	$55,\!10$		10,92	0,001	
Pepsinogen II	$3,\!44$	$1,85 \ 3,80$	$0,\!40$	30,00	$0,90\ 2,40$	$4,\!65\ 5,\!55$	$\hat{a}??$ "p	<
(ng/ml)	$5,\!55$		1,80	36,70			0,001	
?-1-	$1,\!30$	$1,29\ 1,33$	$0,\!48$	$2,\!66\ 2,\!95$	$0,97\ 1,22$	$1,\!62\ 1,\!40$	p > 0.10	
$\begin{array}{c} \text{antitripsin} \\ \text{(g/l)} \end{array}$	1,33		0,84					

3

[Note: Note: ALP -alkaline phosphatase. (numerator -mature newborns, denominator -premature newborns)]

Figure 3: Table 3 :

$\mathbf{4}$

Enzymes	Average	Median value	Minimum	Maximum	Lower quartile	Upper quartile	Shift direc- tion, Statis- tical signifi- cance
Lipase	2,00	$1,\!90$	$0,\!90$	4,90	$1,\!45$	$2,\!35$?p<
(U/l)	4,64	4,05	0,90	33,40	$2,\!10$	4,64	0,01
Amylase	182,73	146,50	37,00	$537,\!00$	92,00	229,00	â??"p<
(U/l)	$67,\!40$	65,70	16,00	162,00	47,00	$84,\!50$	0,01
ALP	$182,\!94$	132,50	18,00	999,00	74,00	$167,\!50$	â??"p<
(U/l)	38,74	$23,\!50$	0,00	$396,\!00$	14,00	44,00	0,01
Pepsinog en	$19,\!28$	$17,\!90$	$7,\!00$	$63,\!10$	$13,\!40$	$19,\!90$	$\hat{a}??"p < 0.01$
I (ng/ml)	33,95	29,85	10,40	$106,00 \\ 1635,9$	24,85	38,00	
Pepsinog en	254,47	124,40	$14,\!10$	$0\ 1631,6$	$57,\!15$	254,47	â??"p< 0,01
II (ng/ml)	545,79	493,05	100,00	0	320,80	733,70	, ,
?-1- antitrypsi n (g/l)	1,21 4,43	0,30 0,30	$0,30 \\ 0,30$	30,00 30,00	0,30 0,30	0,39 0,30	p < 0.005

[Note: Note: ALP -alkaline phosphatase (numerator -full-term newborns, denominator -premature newborns)]

Figure 4: Table 4 :

Enzymes	Average	Median value	Minimum	n Maximum	Lower quartile	Upper quartile	Shift direc- tion, statis- tical signifi- cance
Lipase	43,68	$10,\!65$	$0,\!30$	270,00	2,5	48,75	p>
(U/l)	40,47	20,50	0,10	244,10	4,45	41,94	0,10
Amylase	278,03	204,00	12,00	1289,00	140,00	340,00	â??"p<
(U/l)	92,03	$92,\!03$	$3,\!00$	237,00	44,00	103,00	0,001
ALP	423,23	70,00	21,00	4988,00	32,00	389,00	â??"p<
(U/l)	55,31	46,00	$3,\!00$	397,00	19,50	$55,\!31$	0,001
Pepsino gen	42,70	42,70	0,00	150, 10	19,80	$58,\!15$	â??"p<
I (ng/ml)	$133,\!74$	$90,\!85$	$4,\!10$	$907,\!50$	44,60	133,75	0,001
Pepsino gen	$573,\!64$	388, 15	0,00	2648,70	$148,\!55$	861,00	â??"p<
II (ng/ml)	$1125,\!03$	1108,96	$100,\!60$	$3087,\!80$	$679,\!25$	$1761,\!10$	0,001
?-1-	$0,\!31\ 0,\!32$	$0,\!300,\!30$	$0,\!30$	$0,\!48\ 0,\!45$	$0,\!300,\!30$	$0,\!31\ 0,\!32$	p > 0,10
antitryps in (g/l)			0,30				

[Note: Note: ALP -alkaline phosphatase (numerator -full-term newborns, denominator -premature newborns)]

Figure 5: Table 5 :

glands followed by fetal zymogenic proteases and ?-amylase. 5. The adequate initial digestive potential of the digestive system of newborns is marked by the content of hydrolases in gastric aspirate; the content of hydrolases in amniotic fluid and umbilical cord blood serum are less informative. 6. It is recommended to characterize the initial digestive potential of newborns by the parallel with the results of determination of several hydrolases zymogens mentioned above that were obtained from gastric aspirate at the end of the delivery as well as in the umbilical cord blood serum and amniotic fluids. 7. In immature gestation, the digestive potential of the digestive

system turns out to be reduced differently in different hydrolase systems, but not in the lipase system. 8. The

determination of the initial digestive potential of the digestive system of newborns is promising for justifying the

239 management of their natural, mixed, and artificial feeding.

²⁴⁰ .1 Gratitude

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