This protocol targets canine rabies in patients with normal immunity at any age, who did not receive rabies vaccine, immunoglobulins (RIG, IGIV), or immune suppression.

DO NOT administer rabies vaccine or immunoglobulin (RIG or IVIG). If so, then use MP 7B.

Hospital day (HD)	Risk	Therapy	Testing	Comments
HD 0	Cardiac arrest	DO NOT administer	For Diagnosis:	Prediction is most accurate based on hospitalization
	(dysautonomia) in 27%	rabies vaccine or	Saliva for PCR	for objective signs (not symptoms). First day =0
	over the first 7 days	immunoglobulin (RIG	Skin biopsy for PCR (or	Vaccination after onset of symptoms has never
		or IVIG)	antigen)	worked and may alter immune responses leading to adverse neurological outcomes.
			Serum for antibody	RIG or IVIG delays development of CSF antibodies,
		Minimize stimulation.	CSF for antibody	essential for survival.
		Rabies causes	(Corneal impressions	Sedation is tapered on HD 8 when vagal function
		extensive sensory and	NOT recommended)	ceases. During taper, consider addition of clonidine or
		motor denervation		dexmedetomidine rather than increases in benzodiazepines or ketamine.
		over 10 days. The	Results of testing can	IMPORTANT: vagus nerve function and risk of arrest
		neurological exam is	be delayed. It may be	may persist past HD 7 in patients receiving favipiravir.
		not useful and can	necessary to sedate or	Haloperidol reduces sensory input and is one of 2
		stimulate a cardiac	treat before diagnosis	drugs published for palliation of rabies.
		arrest. Pupillary exam	is confirmed. Sedation	We DO NOT recommend hurst suppression
		remains useful.	for 7 days is less	We DO NOT recommend burst suppression. Ketamine and amantadine are given as
			dangerous than	neuroprotectants based on quinolinic acid in CSF
		Sedation:	untreated rabies.	(excitotoxin) and original use in the successful protocol. Ketamine is anti-nociceptive and avoids
		If alert and no		altering the pupillary response by opiates.
		<u>dysautonomia:</u>	EEG or BIS monitoring	Barbiturates are immunosuppressive and should be
		haloperidol (5 mg		avoided. Propofol appears safe but may cause isoelectric EEG in rabies.
		hourly x 3 doses, the 5	Rabies mimics include	
		mg daily in divided	NMDAR autoimmune	Ventilate using normal parameters. Rabies patients maintain vascular responsiveness to changes in pCO2.
		doses; 0.1 mg/kg	encephalitis, scorpion	Avoid hypocarbia.
		hourly x 3, the 0.1	sting, elapid snake	Please time tracheostomy between HD 8 and HD 12 to avoid periods of known vasospasm and high risk of
		mg/k daily)	venom, Guillain Barre	dysautonomia in the first 7 hospital days.
		If bradycardia or	syndrome, and	
		tachyarrhythmia:	orofacial seizures.	Antipyretics have no effect in rabies. Ambient temperatures may have major effects on heart rate
		ketamine (0.5-1.0		and blood pressure.
		mg/kg/h) + 1:1		
		midazolam. Titrate		
		sedation to prevent		
		dysautonomia with		
		nursing cares.		

HD 0	Poor immune response in dog rabies. Seven of 25 (28%) were in survival range by 7 days (Figure 1).	Inquire IMMEDIATELY about the possibility of investigational or compassionate use of rabies antivirals, biologics, or gene therapies. These require time for approvals and logistics. If the patient has received BCG vaccination, strongly consider repeat intradermal (ID) BCG vaccination as soon as possible, preferably on the bitten limb. Ribavirin SHOULD NOT be used.	Monitoring: Saliva and serum/CSF are tested twice weekly. This is ESSENTIAL for predicting survival/futility and complications	CSF antibody is necessary for survival. Antibody must be detected by HD 7 (in the absence of experimental therapy) for survival. Chlorhexidine oral care interferes with PCR for rabies virus. Freeze saliva. CSF should be sent for cells and protein (criteria for futility) and lactate if available. Favipiravir (ebola oral dosing regimen) modifies the clinical course of rabies (less denervation) but its bioavailability in the brain is uncertain. Some countries have favipiravir available (China, Russia, Japan). There are theoretical reasons to administer BOTH IFN-beta and favipiravir together: interferons reduce the purine precursor pool to enhance inhibition by favipiravir. The serological response to dog rabies is poor. There is plenty of virus antigen in the skin by HD 0. BCG vaccination may serve as a Th-1 immune adjuvant to existing rabies antigen in skin to cross-stimulate the immune response to rabies. A Th1 response promotes antibody formation in the CSF. BCG is used as an immunological adjuvant in cancer therapy. Ribavrin is immunosuppressive. It is an inosine monophosphate dehydrogenase (IMPDH) inhibitor, similar to mycophenolate.
		Consider administration of interferon-beta.		Rabies is inhibited by type I interferons (IFN-alpha and IFN-beta). IFN-alpha can be neurotoxic, but IFN-beta is used to prevent or minimize demyelination in multiple sclerosis. We have only used Avonex 30 mcg IM once weekly. We have anecdotal evidence for a reduction in salivary viral load after IFN-beta. We have not seen adverse effects.
HD 0-16	Increased intracranial pressure	Prevention: Maintain head of bed elevated 30 degrees; Serum sodium > 140 Fludrocortisone; isotonic saline in IVs	Daily serum sodiums	When measured, this averages 20 cm water, which can be compounded by fluid and electrolyte abnormalities and vasospasm (below)

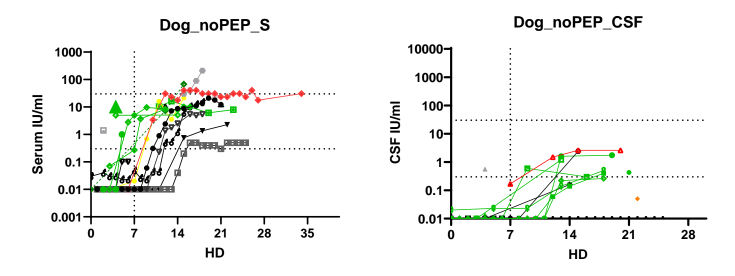
HD 0-16	Low risk of nosocomial transmission	Isolation		There has never been a laboratory-confirmed case of human to human rabies transmission (other than by corneal or solid organ transplantation) during medical care or autopsy. Blood and urine cannot transmit rabies. Patients can be removed from isolation when saliva is negative by PCR on 3 occasions in the presence of neutralizing antibody > 0.5 IU/ml by RFFIT, FAVN or other test for neutralizing antibodies
HD 4	Salt-wasting syndrome on HD 5	Begin fludrocortisone 100 mcg child or 200 mcg adult or other mineralocorticoid; use lactate or normal saline fluids Minimize vasopressors and diuretics; use fluids to maintain blood pressure For hyponatremia, enteric sodium (23%; 1 g in 5 ml water) is more efficacious than 3% IV hypertonic saline	Daily serum sodium and glucose Consider BNP or NT-pro-BNP monitoring Consider serum uric acid and urinary sodium Key test: Saliva for PCR and serum/CSF for neutralizing antibody (HD4)	Avoidance of hyponatremia appears to prevent vasospasm in human rabies. [See HD 6-8] A physiological dose of hydrocortisone (1X dosing) may be used if no pure mineralocorticoid: 15 mg/day divided Q8-12h in adults; 8 mg/m²/day divided Q8h in children. HC risks immunosuppression at higher doses. Central venous pressure appears inaccurate in rabies. Monitoring inferior vena cava collapse by bedside ultrasound is preferred. Rabies is associated with tetrahydrobiopterin (BH4) deficiency that causes adrenaline deficiency as well as nitric oxide (NO) deficiency. Vasopressor agonism is not opposed by NO dilation, leading to profound ileus.
HD 4	Neuroprotection during rabies	Hypothermia is NOT recommended Begin low-dose insulin infusion (0.5 U/h regular insulin in adults; 0.005 U/kg/h in children) and gastric or jejunal feeds.	Daily serum glucose and glucose Urine dipsticks for ketones, daily Begin daily baseline optic nerve sheath diameters by ultrasound (ONSD) or	Hypothermia reduces the immune response This is NOT tight glucose control, rather prevention of catabolism. Reduce insulin if hypoglycemia occurs but maintain some insulin. Complications in rabies are associated with CSF markers of gluconeogenesis and ketogenesis (branched chain amino acids, glycine, acetone, isopropanol) Promotion of anabolism (with adequate caloric intake) appears to improve survival curves by about one week. Insulin may minimize toxic alcohol metabolites and lactic acidosis associated with benzodiazepine sedatives.

HD 4	Thrombosis of cerebral	Amantadine, Vitamin C (500 mg) and vitamin B complex are recommended.	transcranial doppler ultrasounds (TCD) – see cerebral edema	Amantadine (alone with ketamine and midazolam) were part of the original successful protocol. There is biochemical evidence for high quinolinic acid in CSF during rabies (excitotoxicity). Ketamine, midazolam and amantadine are neuroprotectants. Vitamin C recycles BH ₂ to BH ₄ . B vitamins may minimize demyelination.
	veins or sinuses	2 Tr propriy taxes		
HD 5-16	Antibody response causing heart block	Pacer at bedside	Daily ECG HD 5-16	Antibody response to rabies develops between HD 5 and HD 16. Response is very rapid but often is restricted to serum and not CSF. CSF antibody is necessary for survival. Antibody must be detected by HD 7 (in the absence of experimental therapy) for survival. Heart block is associated with the immune response to rabies and progresses over 2-3 days to 3 rd degree block. Pacing works well and blockade recovers over 1-2 weeks. Consider caffeine base (adenosine inhibitor) if no pacer. Atropine no longer effective after 7 days. CAUTION: isoproterenol will dilate intracranial arteries
HD 5-16	Sepsis syndrome			The rabies immune response is associated with increased CRP, WBC with left shift and high platelets. Empirical use of antibiotics should be restricted to 3 days.
HD 6-8	Generalized intracranial vasospasm leading to coma	(fludrocortisone and Na > 140 meq/L)	Optional transcranial doppler ultrasound (needs baseline study) on HD 4-8 and HD 12- 15	Prophylaxis for vasospasm can also be considered using (a) sapropterin (BH ₄ , Kuvan (Merck) (5 mg/kg/day), vitamin C (500 mg) and 0.5 g/kg/day of arginine or citrulline, or (b) nimodipine at ½ to 1/3 the standard dose x 14 days. Sapropterin is preferred but often unavailable. Sapropterin may also improve adrenaline synthesis in the infected adrenal gland. Do NOT use sapropterin and nimodipine together. Vasospasm is associated with onset of coma and mild dysautonomia and pupillary changes. This is evident by transcranial doppler ultrasound and lasts about 1 day, then resolves. Vasospasm is followed by a gradual

HD 7	Survival is associated with detection of neutralizing antibody by HD 7		Key test timing: Serum and CSF for neutralizing antibody (HD 7) Saliva for PCR	increase in intracranial pressure and changes in metabolism (see insulin- above). TCDs are often normal in patients who appear braindead by exam. Lack of detection of neutralizing antibody in CSF by HD 7 indicates medical futility of further care unless you are using an antiviral or other biological. This lab test is essential to unnecessary prolongation of medical care. There is generally a "honeymoon" of medical stability
HD 8	Risk of cardiac arrest abates (unless you use favipiravir or other biological)	Rapidly taper sedation		between HD 8-12 that may be useful for tracheostomy and use of diuretics.
HD 10	Total paralysis by HD 10		Saliva for PCR and serum/CSF for neutralizing antibody (HD 10)	Paralysis reverses (distal to proximal) with virus clearance and often includes orofacial dyskinesias (myokymia) during progression and recovery. These are not seizures.
HD 12-15	Severe generalized intracranial vasospasm; diabetes insipidus		Optional transcranial doppler ultrasound (needs baseline study)	In the absence of neutralizing antibody by HD 7, the patient develops dysautonomia, drop in intracranial pressure and blood flow (high resistance), with flattening of EEG and pupillary dilation, followed in 24 hours by diabetes insipidus.
HD 15 -21	Medical recovery (or criteria for futility)		Saliva for PCR; serum and CSF for rabies neutralizing antibody (HD 15 and weekly thereafter until stable) Criteria for futility (HD >10): diabetes insipidus, isoelectric EEG, CSF lactate > 4 mM, CSF protein > 250 mg/dL	Following "type II" vasospasm, low resistance, chaotic blood flow then supervenes leading to cerebral edema and death. There is no recovery. Virology studies should ALWAYS be completed even if the patient dies. This allows retrospective interpretation of care decisions and the opportunity to detect new complications and improve future care. In the presence of CSF neutralizing antibody (>0.3 IU/ml in our experience), the patient regains pupillary activity, cough and diaphragmatic activity, and reinnervates distal to proximal motor reflexes, then function. Sensation and vestibular function lag. Myokimia re-appears. Dysautonomia occurs with cares and must be tolerated. There are a few days of profuse sialorrhea, bronchial and gastric secretions. There is a subtle SIADH. A few patients develop profound cachexia before recovering. The patients do not regulate temperature well and heart rate (and

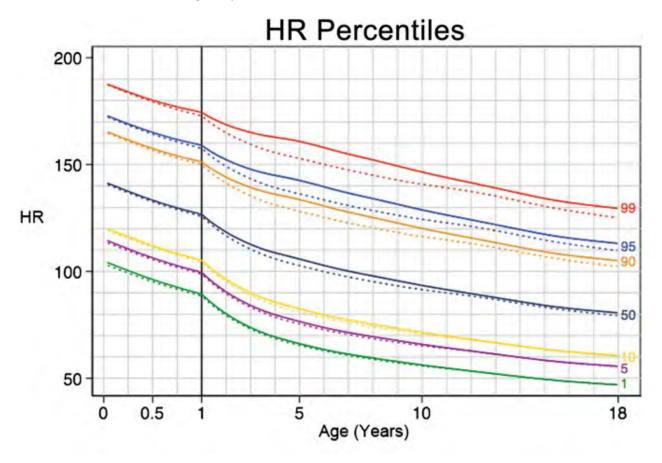
		blood pressure) varies by body temperature; the
		patients require bundling.
Autopsy		There are many ICU complications that result in death during rabies care. The autopsy will identify new complications in 25% of patients. It may show virus clearance (evidenced by lack of virus cultivation and spotty rather than homogeneous detection of virus antigen and RNA in tissues). This finding of virus clearance is often of consolation to family members and the medical staff.
		There are needle biopsy alternatives to standard autopsy when the standard form is prohibited by cultural or religious norms. THERE HAS NEVER BEEN TRANSMISSION OF RABIES DURING AN AUTOPSY.

Figure 1. Immune response to dog rabies without immunomodulation (vaccine, RIG, immunocompromise) in serum and CSF after partial post-exposure prophylaxis, vaccine failure, or vaccination when rabies is symptomatic from our series and the literature. In serum, 28% were in range by 7 days. In CSF, 7% were in range by 7 days.



APPENDIX

Heart rate norms. There is always some tachycardia and fluctuation in rabies. We must tolerate some. By significant dysautonomia that we need to treat with sedation, we refer to heart rates and blood pressures above or below the 99^{th} percentiles for age or height (e.g. P >150 or < 60; BPsys>120 or < 75 in children. For adults, we refer to heart rates and blood pressures above or below the 95^{th} percentiles -- for lack of more extreme normative data (e.g. BPsys > 152 or < 100.



Blood pressure norms (adults)

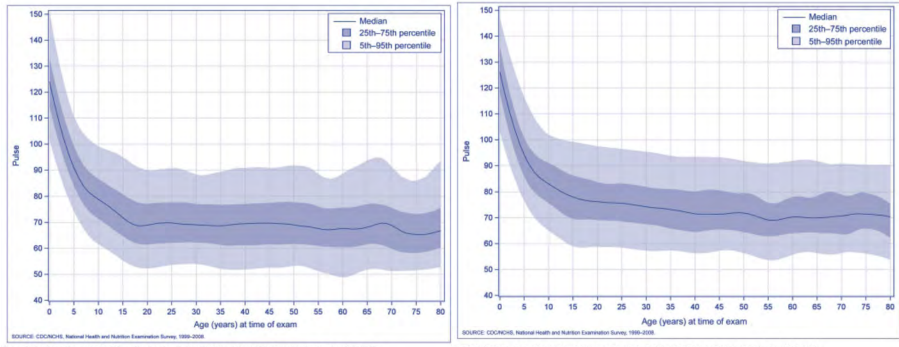
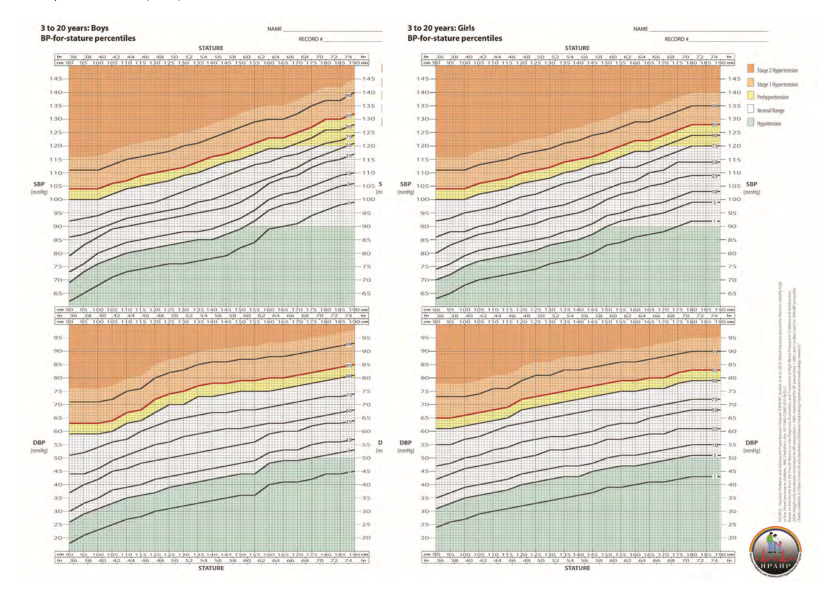


Figure 1. Resting pulse rates for U.S. males, by age: National Health and Nutrition Examination Survey, 1999-2008

Figure 2. Resting pulse rates for females, by age: National Health and Nutrition Examination Survey, 1999–2008

Blood pressue norms (child)



Optic nerve sheath diameter norms (adult)

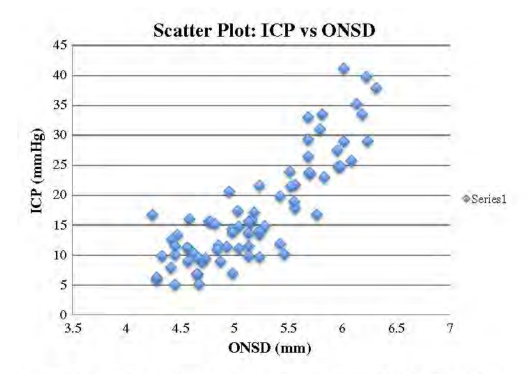


Fig. 1. Scatter Plot 1: ICP vs ONSD. Scatter plot of 75 measurements of ICP in the X axis against the ONSD value in the Y axis. Generally this scatter plot shows a linear relationship. However towards the extreme end of ICP value, the ONSD value started to reach a plateau phase. This is due to maximal dilatation of the optic nerve sheath despite elevation of ICP. Prior studies suggested that with increasing ICPs there might be a maximum nerve sheath diameter that would create an asymptotic relationship. A scatterplot of ICP as a function of ONSD demonstrates this relationship with the maximum ONSD in this population of 6.31 mm.

Optic nerve sheath diameter norms (child)

Table 6 ONSD cut-off values in children >1 year old and children with a closed AF

ICP threshold (in mmHg)	ONSD cut-off in children over 1 year old (in mm)	ONSD cut-off in children with a closed AF (in mm)
<u>≥</u> 20	5.75	5.81
≥15	5.49	5.50
≥20 ≥15 ≥10 ≥5	5.20	5.20
≥5	5.10	5.00

Transcranial doppler ultrasound norms (child): mean flow (time-averaged mean max) and resistive index, middle cerebral artery

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Table 3 Mean (SD) flow velocities in basal cerebral arteries (in cm/second) in a cross sectional study of healthy children

Age	n	Middle cerebral	Internal carotid	Anterior cerebral	Posterior ce artery	rebral	Basilar artery
		artery	artery	artery	P1*	P2†	
Systolic peak flow velocity:							
0-10 days	18	46 (10)	47 (9)	35 (8)		-	_
11-90 days	14	75 (15)	77 (19)	58 (15)	_	_	_
3-11-9 months	13	114 (20)	104 (12)	77 (15)			- Z2
1-2.9 years	9	124 (10)	118 (24)	81 (19)	67 (18)	69 (9)	71 (6)
3-5-9 years	18	147 (17)	144 (19)	104 (22)	84 (20)	81 (16)	88 (9)
6-9-9 years	20	143 (13)	140 (14)	100 (20)	82 (11)	75 (10)	85 (17)
10-18 years	20	129 (17)	125 (18)	92 (19)	75 (16)	66 (10)	68 (11)
Mean flow velocity‡:				750 864			
0-10 days	18	24 (7)	25 (6)	19 (6)	_	_	_
11-90 days	14	42 (10)	43 (12)	33 (11)	-	_	-
3-11.9 months	13	74 (14)	67 (10)	50 (11)			- ·
1-2.9 years	9	85 (10)	81 (8)	55 (13)	50 (17)	50 (12)	51 (6)
3-5-9 years	18	94 (10)	93 (9)	71 (15)	56 (13)	48 (11)	58 (6)
6-9.9 years	20	97 (9)	93 (9)	65 (13)	57 (9)	51 (9)	58 (9)
10-18 years	20	81 (11)	79 (12)	56 (14)	50 (10)	45 (9)	46 (8)
End diastolic peak flow velo	eity:						
0-10 days	18	12 (7)	12 (6)	10 (6)	_	-	_
11-90 days	14	24 (8)	24 (8)	19 (9)	_	·	-
3-11.9 months	13	46 (9)	40 (8)	33 (7)	7	40.00	20.12
1-2-9 years	9	65 (11)	58 (5)	40 (11)	36 (13)	35 (7)	35 (6)
3-5-9 years	18	65 (9)	66 (8)	48 (9)	40 (12)	35 (9)	41 (5)
6-9.9 years	20	72 (9)	68 (10)	51 (10)	42 (7)	38 (7)	44 (8)
10-18 years	20	60 (8)	59 (9)	46 (11)	39 (8)	33 (7)	36 (7)

^{*}Precommunicating part of posterior cerebral artery.

†Postcommunicating part of posterior cerebral artery.

‡Mean flow velocity=time-mean of the maximal velocity envelope curve.

7. Appendices 115

Appendix IVe. Resistance index RI = (vs—vd)/vs—mean values

Age		MCA	ICA	SIPH	ACA	PCA 1	PCA 2	BAS
0-10	days	0.71	0.71*		0.64+	-	-	
11-90	days	0.63	0.71*	-	0.60 +	-		-
3-11.9	months	0.58	0.67*	-	0.60 +	_	_	_
1-2.9	years	0.47	0.52	0.57	0.55	0.55	0.52	0.55
3-5.9	years	0.55	0.60	0.63	0.57	0.58	0.59	0.60
6-9.9	years	0.50	0.55	0.55	0.57	0.55	0.52	0.55
10-16.9	years	0.53	0.58	0.58	0.58	0.55	0.57	0.57

Standard deviations: 0-10 days : 0.11

-11-90 days : 0.07-0.10 3-11.9 months : 0.05-0.07 >1 year : 0.04-0.06

Transcranial doppler ultrasound norms (adult)

TABLE I	200	Reference in Differ			ood Flow V ps	elocitie	es in the Ba	asal C	erebral		
Blood Flow		Subjects									
(cm/sec)	п	Al		20-40	Years Old	41-60	Years Old	>60	Years Old		
ACA	313										
Peak		79 (37	-121)	82	(40-124)	80	(36-124)	72	(52-102)		
Mean		53 (33	-83)	56	(42 - 84)	53	(37-85)	44	(22-66)		
End-diastolic		35 (13	-57)	38	(16-60)	35	(13-57)	28	(12-44)		
MCA	335										
Peak		110 (54	-166)	120	(64-176)	109	(65-175)	92	(58-126)		
Mean		73 (33	-133)	81	(41-121)	73	(35-111)	59	(37-81)		
End-diastolic		49 (21	-77)	55	(29-81)	49	(23-75)	37	(21-53)		
PCA	336										
Peak		71 (39	⊢103)	75	(43-107)	74	(40-108)	62	(38 - 86)		
Mean		49 (25	-73)	52	(28-76)	51	(25-75)	40	(22-58)		
End-diastolic		33 (15	-51)	36	(20-52)	34	(18-50)	26	(14-38)		

ACA = anterior cerebral artery, MCA = middle cerebral artery, PCA = posterior cerebral artery. Range of velocities (calculated as mean ± 2 SD) is given in parentheses.

TABLE 2	Normal Reference Values of Impedance Indexes in the Basal Cerebral Arteries in Different Age Groups							
Impedance		Sub	jects					
Index ^a	All	20-40 Years Old	41-60 Years Old	>60 Years Old				
ACA								
PI	0.87 ± 0.16	0.80 ± 0.14	0.85 ± 0.16	1.02 ± 0.18				
RI	0.56 ± 0.07	0.53 ± 0.05	0.56 ± 0.07	0.62 ± 0.06				
MCA								
PI	0.86 ± 0.15	0.83 ± 0.14	0.82 ± 0.13	0.96 ± 0.17				
RI	0.56 ± 0.06	0.54 ± 0.05	0.55 ± 0.05	0.60 ± 0.06				
PCA								
PI	0.81 ± 0.15	0.76 ± 0.12	0.79 ± 0.12	0.94 ± 0.16				
RI	0.54 ± 0.07	0.52 ± 0.06	0.53 ± 0.05	0.60 ± 0.09				

ACA = anterior cerebral artery, PI = pulsatility index, RI = resistivity index, MCA = middle cerebral artery, PCA = posterior cerebral artery.

a Mean ± 2 SD.