From Neonatal to Fetal Neurology:
Some Clues for Interpreting Fetal Findings

Claudine Amiel-Tison, Julie Gosselin

Correspondence: Claudine Amiel-Tison, Professor Emerita in Pediatrics, Paris V University
Saint-Vincent de Paul Hospital, 82 Avenue Denfert-Rochereau, 75674 Paris Cedex 14, France
Phone: 011 33 1 58412045, Fax: 011 33 1 43 26 12 50, e-mail: amiel-tison@club-internet.fr

Abstract: As early as possible, neonatologists try to identify neonates at risk of unfavorable neurodevelopmental outcomes. They are fairly reliable in predicting very poor outcomes as well as optimal outcomes. However, within these two extremes, the prediction still remains a challenge. Immaturity of the neonatal brain constitutes a limit in itself. During decades with the growing knowledge of brain development, many methods have been developed for neurological assessment of the neonate. Neither of them applied alone was perfect in terms of clinical applicability, sensitivity, reproducibility and specificity.

The motor function is the first to provide the clinician with clues. Higher functions, in particular language and other cognitive functions, will develop later. However, recent researchers give credit to the brainstem for controlling exceedingly rudimentary learning-related cognitive-like activity. At present, the anticipation of late emerging developmental disabilities remains difficult even though early motor dysfunction has repeatedly been associated with a higher risk of intellectual or other learning disabilities. Despite our modest recent contribution to the domain of prediction, further studies on well-defined high risk populations with rigorous methodology that aim to demonstrate these links are still needed. Besides neurological observations, research is in process of including behavioral and stress/reactivity measures; feasibility and benefits have to be demonstrated.

At present, fetal neurology is supported by neonatal neurology. Obstetricians are wise enough to take from both methods described above the elements they are able to transpose to fetal life. A comparative table of neonatal and fetal assessment is to be found elsewhere. As for neonatal neurology, the future of fetal neurology will have to rely on short- and long-term follow-up studies to define the predictive value of the chosen items. Obstetricians will have to be as patient as pediatricians, to work, step by step, towards defining optimality and impairment. They will have to be very careful when deciding to interrupt pregnancies; at the time being, such decisions are restricted to cases of very severe impairment. In line with the spectrum described above, they can expect to find more cases with moderate to mild abnormalities than cases with severe ones. However the most pleasant aspect for the echographer is to check fetal optimality. Just as a newborn infant categorized as at risk of brain damage is competent enough to demonstrate CNS integrity from birth, a high risk fetus will soon be competent enough to demonstrate CNS integrity before birth.

Keywords: Fetal neurology, neonatal neurology, neurological assessment, brain development, brain injury, developmental disability.

OBSTETRICIANS AND NEONATOLOGISTS: A COMMON GOAL

As early as possible, neonatologists try to identify neonates at risk of unfavorable neurodevelopmental outcomes. They are fairly reliable in predicting very poor outcomes as well as optimal outcomes. However, within these two extremes, the prediction still remains a challenge. As the neuromotor system matures early and cranial expansion passively follows hemispheric growth, neurological assessment should be able to produce early markers to predict later outcomes based on neuroromotor and cranial findings. For many years, obstetricians have worked toward the same objective by monitoring fetal well-being during pregnancy. They rely on technical advances, namely ultrasonography (US) which has lead to the following statement: “Fetal behavior can be defined as fetal activities observed or recorded with ultrasound equipment.” [66, p 413] This definition recalls Alfred Binet’s famous answer on being asked what intelligence was: “It is what my test measures”!

In fact, the advent of US has led to a kind of revolution. One of us (CAT) cannot forget the emotion felt when observing a 2D-US fetal assessment and being offered by Jason Birnholz, a pioneer in obstetrical US, a picture of the perfect profile of a 5 month-old fetus named Patricia; it seems one would be able to recognize her 20 years later, based on her facial morphology and expression. At the same time, Hans Prechtl began to describe fetal movements with 2D-US, he used to say that he would be able to recognize years later an individual’s gait pattern based on the quality of his/her fetal movements assessed by 2D-US.

The advent of 3D-4D US constitutes a turning point in the exploration of intrauterine life. In 2005, one of us (CAT) was offered a private session on a 3D-4D US fetal assessment
recorded in the last trimester of pregnancy by Asim Kurjak.45 Looking at this fetus moving was another exciting experience as such an observation shakes our own perceptions of intrauterine life. The quality of movement as well as the variety of facial expressions were striking: a calm fetus, with open eyes and an alert attitude, executing elegant finger and thumb movements. At that time, Asim Kurjak was puzzled by the absence of the Moro reflex, and looked for an interpretation for this surprising finding in an otherwise typical fetus. It was quite likely that the absence of the Moro reflex was the consequence of the head being passively maintained by the uterine wall. Actually, the influence of head stability on motor activity recalls observations made by Albert Grenier on neonates more than 30 years ago: by providing support to the neonate’s head, he made more organized movements possible, that were accompanied by the suppression of the Moro reflex and grasping, as well as an increased state of alertness.31 This clinical research has provided insights into our understanding of the interaction between head control and motor activity within the first weeks of postnatal life. The idea that the fetal environment could have an impact on spontaneous motor activity fits in with the present conception of development, which is viewed as “a complex process in which genetically based and environmentally driven processes continuously interact.” [36, p 1182] One of us (JG) further studied Grenier’s original approach by designing a scoring system allowing for the collection of quantitative data.28 This integration of knowledge on pre- and postnatal behavior has evolved towards the attempt to build a shared clinical view on fetal neurology.9,10

BRIDGING RESEARCH AND PRACTICE

Exhaustive descriptions of the current knowledge on fetal brain maturation are now easily accessible to the clinician.22,41,66 Nevertheless, a brief review may be useful to the reader (Fig. 1). The early stage of neural proliferation and neuronal migration ends by 22 to 23 postmenstrual (PM) weeks. Next, several important events take place: axon and dendrite sprouting, synapse formation, glial cell proliferation and myelination. These processes begin between 20 and 30 PM weeks and extend to 40 PM weeks or beyond. Programmed cell death is known to begin very early, as soon as neurons proliferate, most active between 20 to 31 PM weeks, decreasing soon after, to persist at a lower level all life long. Joseph Volpe’s article, entitled “Subplate neurons—Missing link in brain injury of the premature infant?”70 introduced the importance of the subplate to pediatric literature in 1996. The subplate (SP) is a transient structure which emerges as early as 10 PM weeks, with a maximal volume and activity from 20 to 31-32 PM weeks (being thickest at around 29 PM weeks), located below the future cortical plate, in the space where the white matter will later develop. What is the function of this vanishing structure? The subplate is familiarly compared to a “waiting room” for any axonal routing to or from the cortex, used for a few weeks until target cells are ready. During the waiting period however, temporary afferents contribute to generate fetal behavior.42 Since 1996, extensive clinicopathological data have been provided to clinicians.15,17,35,38,54,71,72 As in many other research fields, new imaging techniques have played a prominent role in these advances,14,20,21,24,39,43,53,55,64 demonstrating normal development and various types of brain impairments according to gestational age (GA). Moreover, experimental data has recently provided new insights into the sensorimotor network; as an example, experiments on Drosophila embryos and larvae highlight the importance of sensory input for the development of the motor function.69 The impact of induced mutation eliminating the sensory function in the Drosophila embryonic peripheral nervous system is expressed in different ways according to the stage of development. In the embryo, the impact is mainly on the ratio of forward/backward peristaltic motor patterns; however, normal head swings still allow normal hatching. Later on, in the larva, the motor behaviors are very abnormal with nearly exclusive backward instead of forward movements. Moreover, head swings, which normally disappear at this stage of development, remain present in the senseless larva. These findings suggest that sensory input is essential for proper functional development of motor circuits. Ghosh and Shatz27 have shown that sensory neurons in the human fetus are present as early as 22 PM weeks in the subplate. However, thalamocortical connections will be functional later on, when target neurons are present in the cortical plate.

FETAL BRAIN DAMAGE IN RELATION TO MATURATIVE STAGES: SELECTIVE VULNERABILITY

Maturative processes, as summarized above, take place very quickly. These changes explain why, for instance, the same hypoxic-ischemic (HI) event will have different consequences according to timing: a similar insult will result in a white matter injury (WMI) at 27 PM weeks and in a grey matter injury (GMI) at 39 PM weeks. Moreover, such HI brain damage will have the same result when occurring in utero or after birth; in other words HI will have identical consequences on a 28 PM week fetus or on a 28 PM week preterm newborn. The main types of brain damage can be briefly described as follows.

White Matter Injury

White matter injury (WMI) is the predominant type of injury, occurring in fetal life or the postnatal period in preterm neonates. WMI is the leading cause of cerebral palsy (CP) of perinatal origin. The particular susceptibility of preoligodendrocytes to anoxia is consistent with abnormalities of myelination found at
various degrees in this population. Fetuses and premature neonates share similar pathogenic mechanisms. If HI or materno-fetal infection occurs, the damage can be cystic periventricular leukomalacias (PVLs) or non cystic WMI injuries, focal or diffuse. The first type (cystic) is easily diagnosed by US and conventional MRI (and may therefore be identified in utero) whereas the second type (non cystic) can only be recognized by Diffusion Weighted Imaging (DWI). Recent studies reviewed by Back15 show: i) a spectrum of WMI including both types (cystic and non cystic), non cystic injury being of milder degree, and ii) a decline in the incidence of cystic PVL and a trend toward an increase in non cystic WMI.

Gray Matter Injury

Neuronal damage associated with WMI can now be identified with new imaging methods as reviewed by Volpe.71,72 Quantitative Magnetic Resonance Imaging (MRI) shows reduced volume in cerebral cortical and deep nuclear structures. Cerebellar lesions are also revealed with quantitative MRI.50-52

Fig. 1: Timetable of Brain Maturation up to 40 PM weeks Developmental process schematically represented up to 40 PM weeks (adapted from 22, 41, 66). A dash-line means that the process is beginning or ending. A bold line indicates the most intensive activity.
Subplate Neuron Injury

This transient structure briefly described above shares with the germinative zone (GZ) the denomination of “vanishing structure”. The SP is of the utmost importance for sensorimotor organization. What happens when SP neurons are damaged by HI? Some of the consequences include the degeneration of waiting neurons deprived of their transient target and a lack of functional synaptic connections with the target neurons. If thalamic afferent axons fail to reach the cortex, sensory processing will be disrupted (see above, experimental senseless drosophilia). The cortical organization will be affected as well. Because of the transient nature of this structure, it will disappear anyway, normal or impaired, leaving predominantly a neuronal disorganization only detectable by functional neuroimaging. Such damage is probably the most common substrate for learning disabilities and behavioral problems in very tiny preterm infants.

Anatomical and Clinical Correlations: Emergence of a Shared Spectrum

Correlations between radiological and clinical data are both possible and important. The two approaches support the existence of a spectrum of abnormalities (Fig. 2). The spectrum of perinatal WMI, ranging from severe (cystic PVL) to moderate or mild (diffuse or focal lesions), is now accessible to neuroradiologists. The spectrum is also accessible to clinicians: A 4-level categorization of neurological abnormalities at 2 years corrected age has been developed, going from CP (disabling or non-disabling) to minor neurocranial signs (clustered or isolated). Even though preliminary results based on this clinical spectrum are promising, more research using concomitant DWI and a structured clinical assessment is needed. Probably due to various factors, such as advances in perinatal care, imaging and standardized clinical assessment, a shift from severe damage to milder types of damage in the spectrum can be observed.

One particular situation remains to be elucidated: early fetal damage or postnatal damage in extremely low birth weight (ELBW) (<1000 g) infants may still be missed by clinical and radiological investigations if the lesions are mainly located in the “vanishing structures” (SP and GZ). Experience shows that the rate of disabling CP in this population is not much higher than in the population of very low birth weight (VLBW) (1000-1500 g) infants but a thorough disorganization expressed in the cognitive and behavioral functions is often identified later on. Alterations of the fetal motor function identified by 3D-4D US may provide the missing link in these cases.

IMPACT OF INTRAUTERINE ENVIRONMENT ON FETAL MOTOR ACTIVITY

Head Stability Passively Maintained

Grenier has demonstrated the essential role of head instability on motor behavior during the first post term (PT) weeks of life. In the experimental condition, manual support is given to the neck and spine while the infant’s alertness and attention are solicited by the examiner: an amazing communication state is reached and facial expression becomes more diversified. At the same time, spectacular changes in motor behavior are observed: jerking movements, Moro and grasp reflexes disappear (an event termed “debugging”), flexor tone in the upper limbs decreases and the infant’s hands open. For a short while, this particular state, called the “liberated state”, allows the infant to intentionally reach and grasp an object (Figs 3 to 5). The significance attributed to this ability transiently obtained in an infant as young as 2 PT weeks is of high predictive validity: a positive response brings additional confirmation of the integrity of the upper system.

Why does demonstration of this “liberated state” contribute to our understanding of fetal motor behavior? Because head stability passively maintained by the uterine wall during the second half of pregnancy obviously creates a permanent situation comparable to the head control transiently obtained postnatally during the experiment described above. Moreover, the facial expression observed during the experiment—calm, concentrated, totally involved in reaching the object proposed—bears a very close resemblance to the facial expression and
attitude of the fetus manipulating its umbilical cord on the midline. In addition, the natural and prolonged “liberated state” of the fetus helps to explain the absence of the Moro and grasp reflexes in utero, as routinely observed by 3D-4D US.\textsuperscript{44-46} It is tempting to speculate on these observations:

1. The fetus is very well protected, including against itself: even though the amount of Wharton’s jelly can protect the umbilical vessels, they are more secure if the grasp reflex is somewhat absent due to head stability in utero. The same remark applies to the Moro reflex, which could be a nuisance if repeatedly induced by any postural changes.

2. Postnatally, the young infant will have a much less “comfy” life from birth until the acquisition of head control, around 6 to 8 PT weeks. Physiological hyperexcitability and very active primary reflexes are well-known transient characteristics during the first 3 months of life. When tested in the parent’s presence, emotion peaks when the newborn begins to walk automatically, precompetence reminiscent of the evolution since homo erectus,\textsuperscript{2} even though this archaic performance cannot be considered useful for the newborn infant or for the neonatologist.

**Freedom from Gravity**

Long before the availability of US, Liley\textsuperscript{49} commented on the differences between fetal and postnatal motor behaviors. For example, he explains the ability to roll over early during fetal life but not from birth to 14 to 20 PT weeks as “... a trick which is simple in a state of neutral buoyancy [which] becomes difficult under the new-found tyranny of gravity”\textsuperscript{5}. At the first minute of postnatal life, the most spectacular response ready to counteract gravity is the full righting reaction observed when the neonate is placed in the upright position: tactile stimulation of the sole of the feet provokes an active response in the extensor muscles; body weight is then actively sustained for a few seconds. As developed below, this response depends on the brainstem (lower motor system) and therefore evolves with the subcortical structures in a caudocephalic wave (with a response restricted to the lower limbs around 28 PM weeks and reaching the head by 38-40 PM weeks). During fetal life, verticality is not at all necessary to elicit the activity of the antigravity muscles: the contact of the soles with the uterine wall stimulates the extensor muscles of the axis, whatever the fetal position is. A large repertoire of fetal movements is possible including trunk rotation due to the freedom from gravity. Moreover, the constant vestibular stimulation coming from both the fetus itself and the mother’s movements probably contributes to the early maturation of the vestibular system\textsuperscript{19} (Fig. 6) (The five cranial nerves with nuclei located in the brainstem (VIII to XII) are mature by 34 PM weeks, i.e. earlier than the ones issued from the pons and midbrain).

Motor acquisitions will appear in the first year of life as a consequence mainly of upper motor system maturation. The infant will successively acquire head control, independent sitting and independent walking, in a descending maturative wave. Incidentally, it is remarkable to observe how the motor system is already well prepared for protection of the infant at each new victory over the “tyranny of gravity”. New postural reactions occur, such as the lateral propping reaction to maintain a sitting position and, later, the parachute reaction, ready to attenuate bad falls while learning to walk.\textsuperscript{6,29} However, in the first 6 to 8
PT weeks, the infant seems to be powerless against gravity; the infant is no longer able to move as was possible in utero. Head control will represent a turning point in the expression of motor activity. Such an understanding of these sequential stages will have to be considered in any early assessment of the neuromotor function.

NEONATAL NEUROLOGICAL ASSESSMENTS

Two Schools, Two Methods

Neurological assessment of the neonate has many facets, continuously evolving according to neurophysiological advances. We will briefly describe two approaches, in other words, two schools: one is the French school initiated by André Thomas and Saint-Anne Dargassies in the fifties. The proposed method is based on an assessment of muscle tone and reflexes, a method inherited by one of us (CAT) in the sixties, which was later transmitted to Julie Gosselin. Recent work has focused on the standardization of the assessment. This tendency represents the so-called “classical method”. The other method from the Netherland school, initiated by Prechtl in the eighties, is based on observing the quality of spontaneous movements. This approach promotes a “change of paradigm from the traditional testing of reflexes and responses to an assessment technique that systematizes the observation of the quality of spontaneous movements [...] which represents] a breakthrough in the functional assessment of the young nervous system.” A high degree of parochialism is easily perceptible in the literature; we must choose the church we want to belong to... A short description of both types of assessment will be presented here prior to a demonstration of their common goal: to focus on upper motor control, which changes along with brain maturation.

The Amiel-Tison Assessment at Term: Based on Tone Changes

Physiological Foundations

André-Thomas was fascinated by brainstem activity, which is so conspicuous in the term newborn infant due to the maturational stage. He defined passive and active tone and considered deviance from the expected tone changes according to maturation to be of valuable significance. As a young neonatologist in the early 1960s, one of us (CAT) had the opportunity to train under his protégée, Saint-Anne Dargassies, assessing neonates in Port-Royal-Baudelocque Hospital in Paris. At that time, this researcher was accumulating data on the ascending wave of maturation from 28 to 40 PM weeks. At the same time in Germany, Peiper was publishing an exhaustive description of the performances revealing cerebral functions in the full term newborn. He described a descending wave of maturation of the upper control system occurring in the first years of life.

It was not until the 1980s, when Harvey Sarnat reviewed anatomical and physiological experiments by Lawrence and Kuyppers that pediatricians became fully aware of the clinical significance of their observations. It then became possible to clinically demonstrate the individual development of both upper and lower motor control systems: i) the lower system, consisting of the brainstem and cerebellum, matures early (becoming accessible to clinical assessment at about 28 PM weeks) in an ascending wave and its essential role is to maintain posture against gravity and flexor tone in the limbs; ii) the upper system, consisting of the cerebral hemispheres and basal ganglia, matures later (clinically emerging at 34 PM weeks) and rapidly for the first 2 years in a descending wave and its essential role is to control the lower system, with relaxation of the limbs and control of the antigravity forces, finally allowing erect posture, walking and fine motor skills (Fig. 7). This distinction became even more relevant after

Fig. 6: Fetal Navigation
Fetuses are passively rocked in their mother’s womb: a long cruise followed by sudden stillness when they land. (from ref 11 with permission)
pathological and radiological data had shown that brain damage in the neonate is mainly located in cerebral hemispheres, in the full term infant with hypoxic ischemic encephalopathy (HIE) or in the premature newborn with periventricular leukomalacia (PVL) and other types of WMI. Consequently, the best predictors of impairment are found in responses that depend on the upper control system and not in the responses that depend mainly on brainstem activity.

Fig. 7: Maturation in motor control from fetal life through infancy

The subcortical pathways (lower system or extrapyramidal, A) derive from the brain stem with maturation proceeding upward, starting in the spinal cord. Their essential role is to maintain posture against gravity. The corticospinal pathways (upper system or pyramidal, B) originate in cerebral hemispheres. Their maturation starts later, proceeds downward from the pons to the spinal cord. They are responsible for control of erect posture and for movements of the extremities including fine motor skills. From term onward, corticospinal control takes over, allowing development of mature head control, sitting and walking. The transition period is in blue.
Elaboration of a Clinical Assessment

These pathophysiological considerations have been the driving force for successive modifications of the clinical assessment at term. More emphasis has been placed on signs that depend on the integrity of the upper structures, such as axial tone and alertness, as well as cranial signs linked to the increase in volume of the cerebral hemispheres. The signs depending on brainstem function, such as primitive reflexes and passive tone in limb flexor muscles have been de-emphasized at the neonatal period as they do not provide information about the cerebral hemispheres and basal ganglia. To gain a more precise definition of infant response, a three-point scale has been devised for each item: 0-normal; 1-moderately abnormal; 2-abnormal. The scoring system has been standardized for application beginning with the full term newborn and continuing up to school age in order to have methodological consistency from birth to 6 years.

Evaluation of head growth and identification of cranial signs are an important part of the assessment. A dramatic increase in brain volume occurs in fetal life and the first two years after term birth. Concomitantly, the skull follows the volumetric increase of the cerebral hemispheres mostly by passive adaptation. The relationship between head and brain growth explains why the classical neurological assessment in infancy universally includes measurement of head circumference (HC). However, this crude information which derives from HC measurement remains insufficient to qualify brain growth. Significant information with respect to the integrity of the underlying cerebral hemispheres can be provided by systematic palpation of the main cranial sutures. In severe cases, every suture is involved, with overlapping being perceived as a ridge. In mild and moderate cases, the squamous suture is particularly informative due to its strategic location at the junction of the cranial vault and the cranial base. Being located between the parietal and temporal bones, it can be felt by palpation just above the ear. The overlapping of this suture has been proposed as a marker of moderately decreased brain growth (Fig. 8).

Passive tone is examined by evaluating the amplitude of slow passive movements (i.e. extensibility of muscles) carried out by the observer when the infant is at rest. Passive tone in the limbs evolves upward from hypotonia up to 28 PM weeks to hypertonia, the normal physiological status at term birth. With brainstem maturation the changes are so fast that one has to define hypo or hypertonia with strict reference to the normal finding at a given PM age. Passive tone in the axis is evaluated by comparing amplitude of passive flexion and passive extension. In the typical situation, at any age, flexion exceeds extension; the reverse pattern is abnormal, as well as unlimited flexion and extension.

Active tone refers to active movements of the infant in reaction to certain situations imposed by the examiner. Three responses are elicited by the following items: i) the active global righting reaction in the upright position; ii) active passage of the head forward during the raise to sit maneuver (Fig. 9); iii) active passage of the head backward during the back to lying posture.
maneuver. These three responses permit analysis of the antigravity forces (lower system) and the control exerted on these antigravity forces by the upper system. The correct technique for each of these three manoeuvres is described elsewhere.4

Spontaneous motor activity is scored on a three-level scale according to both quantitative and qualitative aspects as well as symmetry right/left. Our scoring system (Table 1), is issued from a gestalt approach throughout the neurological assessment. Attention is given to hand opening, independent finger movements, and, above all, to active thumb abduction, as this ability at term is possible only if the upper system is intact.

Table 1: Spontaneous motor activity at 40 PM weeks. Scored 0, 1, 2 for right and left side

<table>
<thead>
<tr>
<th>Spontaneous motor activity</th>
<th>Score</th>
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<tr>
<td>Varied, harmonious</td>
<td>0</td>
</tr>
<tr>
<td>Insufficient, stereotyped</td>
<td>1</td>
</tr>
<tr>
<td>Absent or barely present</td>
<td>2</td>
</tr>
<tr>
<td>Asymmetrical (pathological side) R L</td>
<td></td>
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<table>
<thead>
<tr>
<th>Spontaneous thumb abduction</th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Active thumb</td>
<td>0</td>
</tr>
<tr>
<td>Inactive thumb</td>
<td>2</td>
</tr>
<tr>
<td>Fixed thumb in adduction</td>
<td>2</td>
</tr>
<tr>
<td>Asymmetrical (pathological side) R L</td>
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Primary reflexes are measured by examining a few of them, with particular emphasis on non-nutritive sucking, which is systematically evaluated. Palmar grasping, Moro reflex and automatic walking are considered as adjuncts where unilateral damage is suspected or to confirm CNS depression.

Alertness is evaluated by examining visual fixation and pursuit, quality of sucking and social interaction. When abnormal, hypoactivity, hypotonia, and lethargy will indicate whether the CNS depression is mild/moderate or severe. The excellent inter-observer reliability for the final synthesis was confirmed (kappa coefficient= 0.76).23

Clinical Synthesis

Final synthesis following the analytical evaluation is based on the clustering of signs and symptoms. When every response is optimal (score 0), the probability of a favorable outcome is high. This means that hemispheric growth, alertness/communication, motor control and adaptability are typical of a full-term neonate (Table 2).

When responses are not optimal (score 1 or 2), neurological status can be graded into three categories for full term infants and two categories for preterm infants.

In the full-term neonate (Table 3), three grades of severity can be defined at the end of the first week:

- Mild: Abnormalities of tone hypereexcitability, no CNS depression, no seizures.
- Moderate: Abnormalities of tone, signs of CNS depression, rare or isolated seizures (up to two).
- Severe: Repeated seizures and overt CNS depression.

Repeated assessments in the first week of life will differentiate between two profiles: when signs of CNS depression and tone abnormalities are unchanged from assessment to assessment, the clinical profile is defined as unchanging. This particular clinical profile may indicate a prenatal insult. In this case, three neurocranial signs (Fig. 10) monitored within the first days of life represent essential and specific clues when dating brain damage as prenatal in a full term neonate: a high-arched palate (the result of insufficient tongue movements during fetal life), non-reducible adduction of thumbs in a tightly clenched fist, and suture ridges.3 These signs are not specific per se; they must be already present at birth if they are to be interpreted as having antenatal origins. A recent study confirmed the value of these signs in assessing the etiology and timing of brain lesions.58

| Table 2: Clinical criteria defining optimality of CNS function at term |
|----------------------------------------------------------|-----------|
| Observations or tests                                      | Optimal responses | Significance          |
| Head circumference cm                                      | Same range as birth weight (±25 centiles)   | X  Adequate hemispheric growth |
| Cranial sutures                                            | Edge to edge (squamous included)            | X  |
| Visual pursuit (Fix and track)                             | Easily obtainable                             | X  No CNS depression          |
| Social interaction                                         | Eager                                            | X |
| Sucking reflex                                             | Efficient, rhythmic                            | X  |
| Raise-to-sit and reverse                                   | Active flexor muscles (balance with extensor muscles) | X  |
| Passive axial tone                                         | More flexion than extension                    | X  Upper motor control integrity |
| Passive tone in limbs                                      | Symmetrical and within normal limits for GA    | X  |
| Fingers and thumbs                                         | Independent movements and abduction of thumbs  | X  |
| Autonomic control during assessment                        | Stable color, heart rate, respiration, etc.    | X  Abnormalities suggest ANS* problems |

*ANS: Autonomic Nervous System
On the other hand, when signs of CNS depression increase up until day 3 and then tend to get better with progressive improvements in alertness, motor activity and sucking, the clinical profile is defined as changing. This changing profile is typical of a very recent insult, most often intrapartum.

In the preterm infant, when neurological and/or cranial signs are found around 40 PM weeks, conclusions should take into account extra-neurological problems not yet completely resolved. The degree of severity will be limited to two levels: mild/moderate and severe.

### The Prechtl Assessment, Based on Spontaneous Motor Behavior

**Physiological Foundations**

Prechtl and his followers proposed a conception of motor behavior which is understood as “the net result of the activity of complex spinal or brainstem machineries, which are subtly modulated by segmental afferent information and ingeniously controlled by supraspinal networks” (p 483). In other words, the young nervous system generates a variety of motor patterns originating in “neural networks that are able to coordinate autonomously (i.e. without sequential sensory or supraspinal information) the activity of many muscles” (p 483). The observation of pre- and postnatal spontaneous motor behavior drove Prechtl to describe general movements (GMs) as “series of gross movements of variable speed and amplitude, which involve all parts of the body but lack a distinctive sequencing of the participating body parts” (p 484).

Due to the timing of neuronal maturation and the occurrence of GMs, this motor pattern is considered as being produced without afferent information. “Remarkably, GMs are among the first movements that the human fetus develops, and they emerge prior to isolated limb movements. GMs can already be observed before the completion of the spinal reflex arc, which is accomplished at 8 weeks postmenstrual age (PMA)” (p 484).
and neuromaturative phenomena. These correlations are mainly based on the “coincidence of emergence”:

- During the third trimester of pregnancy: Improvement in movement complexity and variation and the emergence of synaptic activity in the subplate;
- Near term: Emergence of “writhing GMs” and changes in cortical neurotransmitter systems.

Elaboration of a Clinical Assessment

Prechtl’s initial clinical studies came with the advent of advanced ultrasound equipment which made possible prolonged and repeated direct observations from 7 to 8 PM weeks onwards. Continuous one-hour recordings are made and stored on videotape for off-line analysis. As for the ex utero examination, a video camera is installed high above the infant who is lying supine in the incubator or bed. Recordings are made while the infant is in state of alertness 3 and last between 30 and 60 minutes in order to collect about three GMs. Later on, recordings are viewed at high speed and GMs are then appraised on a visual gestalt perception of the spatial and temporal variation of motor patterns. The interjudge and intrajudge reliabilities are considered to be excellent, with Kappa values around 0.80 achieved after 2 days on site-training and further practice of about 100 GM recordings.

The in utero assessment has permitted a description of the fetal motor repertoire, and demonstrates the influence of GMs on changes in fetal posture: the contact of the feet on the uterine wall resulting from GMs and alternating leg movements provoke changes in the intrauterine position in utero. From 36-38 PM weeks until the end of the second month post-term, the repertoire changes and GMs are commonly referred to as writhing movements, characterized by small to moderate amplitude and slow to moderate speed. They are ellipsoid in form, which creates the impression of a writhing quality. By 2 months post term, they gradually disappear and fidgety GMs gradually emerge: small movements in all directions of moderate speed and variable acceleration involving the neck, trunk and limbs; they last until 20 PT weeks. A temporal overlap can be observed as shown in Table 4.

Table 4: Age-specific characteristics of normal GMs (from ref 35, with permission)

<table>
<thead>
<tr>
<th>GM type</th>
<th>Period of presence (weeks PMA)</th>
<th>Description</th>
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<tbody>
<tr>
<td>Preterm GMs</td>
<td>From ± 28 w until 36-38 w</td>
<td>Extremely variable movements, including many pelvic tilts and trunk movements</td>
</tr>
<tr>
<td>Writhing GMs</td>
<td>Form 36-38 w until 46-52 w</td>
<td>The variable movements take on a more forceful (“writhing”) character. In comparison with preterm GMs, “writhing” GMs seem to be somewhat slower and to show less participation of the pelvis and trunk</td>
</tr>
<tr>
<td>Fidgety GMs</td>
<td>From 46-52 w until 54-58 w</td>
<td>Basic motility consists of a continuous flow of small and elegant movements occurring irregularly all over the body, i.e. head, trunk, and limbs participate to a similar extent. The small movements can be superimposed on large and fast movements.</td>
</tr>
</tbody>
</table>

At any GM age, the basic characteristics of normal GMs are: (i) participation of all body parts and (ii) movement complexity and variation.
Clinical Synthesis

If the nervous system is impaired, GMs lose their complex and variable character. Different types of abnormal patterns may be recognized and clustered into different age-specific categories:

- **Poor repertoire** when the sequence of the successive movement components is monotonous and movements of the different body parts do not occur in the complex way seen in normal GMs.
- **Cramped-synchronized GMs** are considered to be abnormal from preterm age onwards. These cramped-synchronized GMs refer to rigid movements lacking the normal smooth and fluent character; all limb and trunk muscles contract and relax almost simultaneously. If this abnormal pattern is observed consistently over a few of weeks it is of high predictive value for the development of spastic CP.
- **Chaotic GMs** of all limbs are of large amplitude and occur in a chaotic order without any fluency or smoothness. Infants with chaotic GMs often develop cramped-synchronized GMs a few weeks later.
- **Abnormal fidgety movements** look like normal fidgety movements but their amplitude, speed and jerkiness are moderately or greatly exaggerated.
- **Absence of fidgety movements** is concluded when movements are never observed from 9 to 20 PT weeks; this absence is highly predictive for later neurological impairment. Predictive value at both extremes of the spectrum is good: cramped synchronized GMs are highly predictive of CP; normal repertoire is highly predictive of an optimal outcome. However, in between, the group of infants with a poor repertoire will later split into normal, subnormal or severely damaged subgroups.

Similarities and Differences

In the upper section of Table 5, similarities are analyzed. In the lower section, differences are highlighted. As far as reliability and predictive value are concerned, both methods are comparable. However, both methods face the same challenge in intermediate situations, for which outcomes remain difficult to establish before a more advanced level of neuromaturation. As for the method itself, the “hands off” observation of GMs as proposed by Prechtl has an obviously invaluable advantage for fetal 3D-4D observations as well as for very sick infants. On the other hand, such a distance between the examiner and the infant may represent a significant source of frustration for pediatricians. The latter may be more comfortable with the “hands on” situation proposed in the classical examination, which creates interaction between examiner and examinee. The infant’s behavior is relational, as proposed by Neisser; continued participation and mutuality between the neonate and the examiner is experienced daily by “hands on” clinicians (Fig. 11). Moreover, such contact may allow for a better appraisal of the different alertness states as well as the fluidity of changes. Finally, encompassing the first 6 years of life, the classical assessment provides methodological continuity, continuity of the utmost importance in the course of any follow-up.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>ATNAT</th>
<th>GMA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration of assessment</td>
<td>Comparable</td>
<td></td>
</tr>
<tr>
<td>Interobserver reliability</td>
<td>Similar</td>
<td></td>
</tr>
<tr>
<td>Predictive value</td>
<td>Similar</td>
<td></td>
</tr>
<tr>
<td>For disabling CP</td>
<td>Good</td>
<td></td>
</tr>
<tr>
<td>For non-disabling CP</td>
<td>Poor</td>
<td></td>
</tr>
<tr>
<td>For optimality</td>
<td>Good</td>
<td></td>
</tr>
<tr>
<td>Identification of prenatal damage</td>
<td>Comparable</td>
<td></td>
</tr>
<tr>
<td>Type of assessment</td>
<td>Hands-on</td>
<td>Hands-off</td>
</tr>
<tr>
<td>Feto-neonatal continuity</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Continuity in the course of follow-up</td>
<td>Yes (6 years)</td>
<td>No</td>
</tr>
</tbody>
</table>

Fig. 11: A 2 days old baby girl
She is self quieting at the end of the assessment: peaceful recovery, holding on the midline and sucking the examiner’s finger.

The two methods differ as regards with the specific manoeuvres or signs used to test the integrity of the different neuro-pathways (Fig. 12). Spinal and brainstem control are omnipotent until 34 PM weeks or so. A transitional period follows during which corticospinal control progressively becomes preponderant. Head control is a very important marker, accompanied by the decrease of primary reflexes and smoother motor activity, making life more comfortable around 8 to 10 PT weeks. Later on, the corticospinal power over lower structures will advance, allowing gross and fine motor acquisitions. However, both methods explore subcorticospinal as well as corticospinal control, both recognize a transitional
phase from around 34-36 PM weeks to 6-8 PT weeks: the GMA explores this transition through writhing movements while the ATNAT mainly uses the active tone in the flexor muscles of the body axis and active abduction of the thumbs.

**EXPECTATIONS AND LIMITS OF FETAL NEUROLOGY**

**Expectations**

Throughout this chapter devoted to methodology in the rapidly changing field of neonatal neurology, we have tried to combine experience and reason. We have chosen to compare our own assessment, belonging to the so-called “classical” school, with the GMA, in which we have little personal experience. We have therefore tried to stay as close as possible to the original writings, hence our quotations taken from the texts of Prechtl and other colleagues. During our readings, we have noticed the stereotypical way in which the classical assessment is labelled “reflex neurology”, based on limited references confined to André-Thomas and Peiper taken from the early sixties. Those two researchers were great clinicians but did not have access to the amazing recent advances in neurosciences. Moreover, their research on automatic walking is still current, as shown by the number of publications concerning this fascinating primary locomotion. For instance, recent experiments argue in favor of a rudimentary coupling present at birth between optical flow and locomotion. Such findings demonstrate that the infant’s behavior is differentiated in line with the task at hand.

Throughout these methodological comparisons we have tried to establish links between the two approaches; we were struck by the fact that the goal of each approach is exactly the same and deals with the development of early markers indicating...
impairment of the upper structures of the brain. In this respect, current data show that any type of assessment is good enough to define optimality on the one hand and severe impairment on the other hand (as well as mothers, caregivers … or radiologists with fetal IRM!). In between these two extremes, they face the same challenge, with the risk that early intervention in children who will later suffer from developmental disabilities is postponed.

Limits

Immaturity of the neonatal brain constitutes a limit in itself. The motor function, being the first to mature, is the first to provide the clinician with clues. Higher functions, in particular language and other cognitive functions, will develop later. However, recent researchers give credit to the brainstem for controlling exceedingly rudimentary learning-related cognitive-like activity. At present, the anticipation of late emerging developmental disabilities remains difficult even though early motor dysfunction has repeatedly been associated with a higher risk of intellectual deficiency or other learning disabilities. Despite our modest recent contribution to the domain of prediction, further studies on well-defined high risk populations with rigorous methodology that aim to demonstrate these links are still needed. Besides neurological observations research is in process of including behavioral and stress/reactivity measures; feasibility and benefits have to be demonstrated.

CONCLUSION

At present, fetal neurology is supported by neonatal neurology. Obstetricians are wise enough to take from both methods described above the elements they are able to transpose to fetal life. A comparative table of neonatal and fetal assessment is to be found elsewhere in this book. As for neonatal neurology, the future of fetal neurology will have to rely on short- and long-term follow-up studies to define the predictive value of the chosen items. Obstetricians will have to be as patient as pediatricians, to work, step by step, towards defining optimality and impairment. They will have to be very careful when deciding to interrupt pregnancies; at the time being, such decisions are restricted to cases of very severe impairment. In line with the spectrum described above, they can expect to find more cases with moderate to mild abnormalities than cases with severe ones. However the most pleasant aspect for the echographer is to check fetal optimality as beautifully illustrated in various texts by Kurjak’s school. Just as a newborn infant categorized as at risk of brain damage is competent enough to demonstrate CNS integrity from birth, a high risk fetus will soon be competent enough to demonstrate CNS integrity before birth.

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