Comparable Measures of Cognitive Function in Human Infants and Laboratory Animals to Identify Environmental Health Risks to Children

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The importance of including neurodevelopmental endpoints in environmental studies is clear. A validated measure of cognitive function in human infants that also has a homologous or parallel test in laboratory animal studies will provide a valuable approach for large-scale studies. Such a comparable test will allow researchers to observe the effect of environmental neurotoxicants in animals and relate those findings to humans. In this article, we present the results of a review of post-1990, peer-reviewed literature and current research examining measures of cognitive function that can be applied to both human infants (0–12 months old) and laboratory animals. We begin with a discussion of the definition of cognitive function and important considerations in cross-species research. We then describe identified comparable measures, providing a description of the test in human infants and animal subjects. Available information on test reliability, validity, and population norms, as well as test limitations and constraints, is also presented. Key words: attention, behavioral testing methodology, cognitive function, developmental neurotoxicology, environmental health, infant, intelligence, learning, memory, neurobehavior. Environ Health Perspect 111:1630–1639 (2003). doi:10.1289/ehp.6205 available via http://dx.doi.org/[Online 2 July 2003]

Impairment of cognitive function is a recognized primary outcome of exposure to developmental neurotoxicants, such as lead, methyl mercury, polychlorinated biphenyls (PCBs), and other chemicals. Efficient inclusion of this end point in environmental studies will rely on a validated measure of cognitive function in human infants that has a parallel test in laboratory animal studies. The identification of a comparable measure of cognitive function in human infant and animal studies will facilitate toxicology studies designed to evaluate mechanistic and dose–response aspects of effects observed in human infants.

In this article, we present the results of a review of post-1990, peer-reviewed literature examining measures of cognitive function that can be applied to both human infants (0–12 months old) and laboratory animals.

What Is Cognitive Function?

“Cognition” is vaguely defined as “the act or process of knowing, including both awareness and judgement” (Merriam-Webster On-Line: The Language Center 2003). Hence, it is important to define cognitive function in the context in which it is used. For this article, we define “cognitive function” as encompassing learning, memory, and attention processes (Cory-Slechta et al. 2001). “Learning” is classically defined as a relatively permanent behavior change as a result of practice or experience. When an infant or young animal responds in an adaptive way to a stimulus, learning (or information processing) has occurred (Fagen and Ohr 2001). “Memory” is then defined as the persistence of a learned behavior over time (U.S. EPA 1998). “Attention” refers to a global behavioral construct that includes numerous response classes such as impulsivity, sensitivity to delay, activity level, sustained attention, and ability to manage delay of reward (Bushnell 1998; Bushnell and Rice 1999; Cory-Slechta et al. 2001). In infants, attention research has focused on four areas of visual attention: alertness, spatial orienting, attention to object features, and endogenous or internally directed, attentional functions (e.g., attention span, perseverance, and distractibility; Colombo 2001).

Cross-Species Developmental Neurotoxicity

The adverse effects of developmental exposure to neurotoxicants on various cognitive functions can be assessed in both humans and animals. However, the degree to which specific assessment techniques are comparable across species can vary dramatically. The 1990 Workshop on Qualitative and Quantitative Comparability of Human and Animal Developmental Neurotoxicity (Stanton and Spear 1990), sponsored by the U.S. Environmental Protection Agency and the National Institute on Drug Abuse, proposed four criteria for evaluation of such animal models: a) Developmental profiles of functional capacity should resemble those found in humans; b) conceptual or operational similarities should exist between behavioral measures of those capacities in developing humans and animals; c) developmental profiles of neurobiologic changes should resemble those found in humans, particularly those that underlie the functional capacity in question; and d) treatments that alter neural or behavioral maturational in humans should cause similar alterations in the animal model.

Over the past decade, neurotoxicologists have directed considerable effort toward modeling human cognitive function in animals and applying animal cognitive function tests to humans (Adams et al. 2000; Anderson 2000; Paule 2001; Paule and Barone 2000). Examples include the following: a) The operant battery test (OTB) from the U.S. Food and Drug Administration’s National Center for Toxicological Research (NCTR), used with laboratory rhesus monkeys, has been successfully applied to assessments in 6-year-olds (Paule et al. 1999a, 1999b; Slikker et al. 2000). Performance of children on money reinforcement (nickels) operant tests of motivation, color and position discrimination, learning, short-term memory, and time estimation were compared with standardized IQ (intelligence quotient) tests. Many tests in the OTB have also been adapted for use in rats (Mayorga et al. 2000a, 2000b), b) The Wisconsin General Testing Apparatus, radial-arm maze, and the Morris search apparatus, used to test cognitive function in nonhuman primates or rodents, have been successfully adapted for tests of toddlers and preschool children (Overman 1990; Overman and Bachevalier 2001; Overman et al. 1996a, 1996b). c) The well-studied Computer-Assisted Neurotoxicology Assessment Battery, developed for older children and adults, has also been applied to animal models (Fray and Robbins 1996).

The models presented above have not been applied in infants. Similar applications of tests in animals to the study of human infants present obvious obstacles. Human infants lack language, display poorly developed motor skills, and undergo a prolonged period of infancy.

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Nevertheless, a wide body of research in developmental psychology shows that infants, even newborns, learn, remember, and focus attention (Nelson and Luciana 2001). For risk assessment, the neurobehavioral assessment of infants presents two challenges (Bellinger 2002). First, the highly dynamic nature of early neurodevelopment presents a moving target, making it difficult to interpret apparent performance deficits in the absence of a baseline measure. Second, normal change over time is expected and must be distinguished from a deviation that may be triggered by neurotoxicant exposure.

Research in human infants has focused mainly on simple forms of learning such as habituation and classical conditioning, where the young infant’s behavior is changed as a function of specific experience, and through which the memory store of the aging child is altered over successive life events (Lipsitt 1990). Operant learning tasks, in which the infant or animal must manipulate a specific part of their environment to receive a reinforcer, are possible only when the infant acquires sufficient motor skills for the task and thus are often limited to age 6 months and older.

**Comparable Measures of Cognitive Function**

Ultimately, neurobehavioral toxicologists seek a sensitive homologous or parallel test in human infants and laboratory animals that can distinguish normal subjects from those that have had an exposure to a neurotoxicant. Although tests of cognitive function can be performed in a variety of species and age groups, this review is limited to studies in rodents, nonhuman primates, and human infants (0–12 months old).

Table 1 presents an overview of tests described here, identified as either homologous or parallel for each species that has been studied. Homologous tests are those for which the same procedure is followed in humans and the animal species. Parallel tests are those that are conducted in a different manner in humans and the animal species, but for which it is believed the same cognitive function is being measured. Table 2 summarizes information for each of the tests.

### Eye-Blink Conditioning

Eye-blink conditioning (EBC) is a model system for studying neural correlates of learning and memory (Sears and Steinmetz 2000; Stanton and Freeman 1994; Woodruff-Pak and Steinmetz 2000a, 2000b). Data collected from human and animals (monkeys, rabbits, rats, cats, mice) show similar patterns of acquisition, retention, and extinction of EBC. Analysis of neural systems and structures involved in EBC have been documented through studies employing stimulation, lesion, and pharmacologic methods. Data collection has consistently demonstrated that brain networks used in EBC are virtually identical across vertebrate species, including humans, monkeys, rabbits, rats, cats, and mice. EBC can be used in the same way for comparison studies across the life span. EBC can distinguish between normative groups and populations with impaired learning or memory disorders, such as between normal and autistic children or between normal aging and Alzheimer disease.

The EBC procedure involves pairing a conditioned stimulus (CS; typically a pure tone) and an unconditioned stimulus (US; typically a brief air puff to the eyelid area). The EBC task can be varied by changing the length of the trace or complexity of the conditioning stimuli, or by methods such as discrimination reversal conditioning (Sears and Steinmetz 2000). There is evidence that delay EBC (when the CS and US overlap and coterminate) can be acquired and retained independently of the forebrain and independently of awareness, whereas trace EBC [which occurs when a short empty interval called the interstimulus interval (ISI) separates the CS and US] cannot (Manns and Clark 2002). In delay EBC, the memory trace is localized in the cerebrum, although the hippocampus is also engaged in the acquisition of a conditioned eye-blink response. Trace EBC depends critically on the cerebellum, but also on the hippocampus if the trace interval is sufficiently long (Kishimoto et al. 2001).

**Infant model.** Although EBC has been well studied in adults, considerably less work has been done in human infants and children. The developmental aspects of the conditioned response have not been systematically studied using either a cross-sectional or a longitudinal approach (Sears and Steinmetz 2000). From limited published data on normal infants and

**Table 1.** Homologous (H) and parallel (P) measures of cognitive function in animals and human infants (X indicates no measure identified).

| Task                                           | Cognitive function assessed                                    | Rodents | Nonhuman primates | Human infants |
|-----------------------------------------------|----------------------------------------------------------------|---------|--------------------|---------------|
| Classical eye-blink conditioning (EBC)        | Associative learning                                           | H       | H                  | H             |
|                                              | Short-term memory                                              |         |                    |               |
|                                              | Attention                                                      |         |                    |               |
|                                              | Inhibitory learning                                            |         |                    |               |
| Visual habituation/novelty preference; visual recognition memory | Visual recognition memory                                      | P       | H                  | H             |
|                                              | Attention to novelty                                            |         |                    |               |
| A-not-B; delay-tolerance A-not-B              | Working memory                                                 |         |                    |               |
|                                              | Spatial memory                                                 |         |                    |               |
|                                              | Inhibitory control                                             |         |                    |               |
| Transparent barrier detour (also called object retrieval) | Working memory                                                 | X       | H                  | H             |
|                                              | Spatial memory                                                 |         |                    |               |
|                                              | Inhibitory control                                             |         |                    |               |
| Mobile/train conjugate reinforcement         | Learning                                                       | P       | H                  | H             |
|                                              | Long-term memory                                               |         |                    |               |
| Delayed nonmatching to sample (DNMS)         | Learning                                                       | X       | H                  | H             |
|                                              | Motivation                                                     |         |                    |               |
|                                              | Working memory                                                 |         |                    |               |
| Means–end problem solving                    | Learning                                                       | P       | H                  | H             |
|                                              | Motivation                                                     |         |                    |               |
|                                              | Memory                                                          |         |                    |               |
| Event-related potentials                     | Recognition memory                                             | P       | H                  | H             |
| Operant discrimination (object features and spatial mapping discrimination) | Learning                                                       | P       | H                  | H             |
|                                              | Memory                                                          |         |                    |               |
|                                              | Attention                                                      |         |                    |               |
| Bayley Scales of Infant Development II (or BSID II) | Number of behavioral and reflex tasks                          | X       | H                  | H             |
Table 2. Summary of comparable measures of cognitive function.

| Task | Summary of test | Equipment required |
|------|-----------------|--------------------|
| **Classical eye-blink conditioning (EBC)** | Pavlovian conditioning procedure involves pairing a conditional stimulus (CS; typically a pure tone) and an unconditional stimulus (US; typically a brief air puff to the eyelid area). The air puff elicits a reflexive eye blink and, after repeated conditioning trials, the response comes to be evoked by the tone CS before or in the absence of the air puff US (Stanton and Freeman 1994). Variations: delay EBC, CS, and US overlap and coterminate; trace EBC, an ISI separates the CS and the US. | Two rooms: one for parents and infant preparation, one for task setup. Standardized visual display of brightly colored objects. Soft band to secure the infant’s head. Flexible plastic tube to deliver air-puff to right eye. Two small 7-ohm speakers to deliver tone CS (1 kHz, 80 dB). Background music. Two cameras to video the infant’s head. Signal box with counter and indicator lights for tone and air puff. EMG recording equipment. |
| **Transparent barrier detour (object retrieval)** | Toy (treat) is placed in box within easy reach of subject. There is a strong pull to reach straight for the toy through the side one is looking, which must be inhibited when subject is looking through closed side of box. | Small clear box in which to place toy or treat, open on one side only. |
| **A-not-B; delay tolerance A-not-B** | The subject (infant or monkey) watches as a reward (toy for infants) is hidden to the left or right in one of two identical locations (A or B). A few seconds later, the subject is encouraged to find the hidden treat. The reward for correct reaching is the toy (or treat). After successful retrieval of the toy (or treat) from location A on two consecutive trials, it is hidden in location B with the subject watching. Measures: A-not-B, correct vs. incorrect location reached on the reversal trial (location B); delay-tolerance A-not-B; Length of longest delay the subject can tolerate and still succeed in retrieving the toy on reversal trials (Diamond 2001a). | Procedural variations: location of ultimate hand motion in hiding sequence, distance between hiding locations, distribution of reaches on warmup trials, differences in covers of background surface, presence of distraction during delay, room illumination, and criterion for determining whether reach is correct (Diamond 2001c; Noland 2001) |
| **Visual habituation/novelty preference; visual recognition memory** | Paired comparison: The infant is presented with a single or two identical targets for a period of familiarization. The familiar target is then paired with a novel one. The extra time spent looking at the novel target implies recognition memory. Nine or 10 comparisons are usually used in a session. | Targets: abstract patterns and shapes (Colombo 1993), or a combination of faces and abstract patterns (Rose et al. 2001b). A three-sided, curtained enclosure with a pivoting stage for presentation of paired stimulus targets. Peephole located midway between the two stimuli for observation of infant corneal reflections of stimulus patterns. Computer for recording looks and looking time and controlling the timing of trials (Rose et al. 2001a). |
| **Classical eye-blink conditioning (EBC)** | Pavlovian conditioning procedure involves pairing a conditional stimulus (CS; typically a pure tone) and an unconditional stimulus (US; typically a brief air puff to the eyelid area). The air puff elicits a reflexive eye blink and, after repeated conditioning trials, the response comes to be evoked by the tone CS before or in the absence of the air puff US (Stanton and Freeman 1994). Variations: delay EBC, CS, and US overlap and coterminate; trace EBC, an ISI separates the CS and the US. | Two rooms: one for parents and infant preparation, one for task setup. Standardized visual display of brightly colored objects. Soft band to secure the infant’s head. Flexible plastic tube to deliver air-puff to right eye. Two small 7-ohm speakers to deliver tone CS (1 kHz, 80 dB). Background music. Two cameras to video the infant’s head. Signal box with counter and indicator lights for tone and air puff. EMG recording equipment. |
| **Transparent barrier detour (object retrieval)** | Toy (treat) is placed in box within easy reach of subject. There is a strong pull to reach straight for the toy through the side one is looking, which must be inhibited when subject is looking through closed side of box. | Small clear box in which to place toy or treat, open on one side only. |

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Means–end problem solving: At 7–8 months, task involves placing a cloth in reach of child and child must remove barrier to grasp cloth, pull cloth to retrieve cover, and search under cover to find toy (two intermediate steps). At 10 months, infants must remove barrier to grasp cloth, pull cloth to retrieve cover, and search under cover to find toy (three intermediate steps). For 10-month-old, infants receive several trials to solve problem. Score is based on criteria for evidence of intention to retrieve the hidden toy (Willatts and Forsyth 2000).

Operant discrimination (object features and spatial mapping discrimination): Visual/spatial displays are presented to the right and left of midline. Looking to a “correct” dimension (color, form, or spatial position) produces synchronous auditory reinforcement. Measures retention of correct dimension.

Event-related potentials (ERPs): Evaluation of a synchronized portion of the QEEG, time-locked to the onset of some event in the infant’s environment. Limitations: The procedure has significant constraints, including problems of between-subject variability in placement of electrodes on the scalp, choice of reference electrode location, and muscle and other forms of artifacts (Marshall and Fox 2001).

Testing scales: Bayley Scales of Infant Development II: Individually administered instrument composed of two main subscales: mental scale, 118 items that assess mental ability (memory, habituation, problem solving, ability to vocalize, language and social skills); motor scale, 111 items that assess motor ability (rolling, crawling and creeping, sitting, standing, walking, running, jumping). All items arranged in order of developmental difficulty. Specification provided for specific sets of items to administer to a child depending on chronological age (Bayley 1993). Early Childhood Longitudinal Study reduced-item Bayley (ECLS-B): A reduced-item set developed that can be administered in less time and produce reliable, valid scores equivalent to the full set (West and Andreassen 2002). Items have been selected for their operational ease and psychometric properties. Multiple items can be scored from one administration, and, in the motor specialty, several items can be scored from observation.

Infants at 3 and 6 months of age are conditioned to move an overhead crib mobile by kicking one of their feet (mobile conjugate reinforcement). At 9 and 12 months, infants are conditioned to activate a musical train and a bank of 10 lights with a lever press response. At each age, 15-min conditioning sessions are conducted in a series of home visits separated by 24 hr. After conditioning sessions, infants are tested after increasing delays (1, 7, or 14 days later) until they exhibit no retention for 2 successive weeks.

Contour detection and closure detection: Using the mobile described above, the infant learns to kick to move the mobile. After two learning sessions on 2 consecutive days, one or more visual characteristics (contour and closure) of the mobile are altered for some infants and not for others. On the third test day, recognition and discrimination of the test mobile are assessed using kick rates in the presence of the training mobile (old) or a novel mobile (new) relative to a baseline acquired for that infant before learning the task.

Delayed nonmatching to sample (DNMS): A sample object is presented. A delay follows, and then the familiar object is presented alongside a novel object. The correct choice is to select the novel object.

Stimulus: treat or toy
Mobile or musical train with lighted press response box
Limitations: Test is labor intensive
Significant respondent burden
Infant motivational factors also impact on test.

Task cannot be automated, so problems associated with tester–subject interaction must be addressed.

Children, Sears and Steinmetz (2000) described the developmental process. Between infancy and early childhood, the acquisition rate for the conditioned eye-blink response dramatically increases from 28% at 1 month to levels near 80% at 5 months, and near 70% for 4- to 6-year-olds. These conditioning rates are similar to rates seen in adults, although the optimal ISI required for conditioning varies from adult protocols. In 5-month-old infants, a delay of 650 msec produces more robust conditioning than do intervals of either 250 or 1,200 msec (Ivkovich et al. 2002).

In the first use of this procedure, 61.5% of 4- and 5-month-olds did not yield reliable data either because they failed to achieve the criterion number of trials (30 tone–air puff trials) or because of technical or procedural problems (Ivkovich et al. 2000). In a later study (Ivkovich et al. 2002), the attrition rate was reduced to 34%. The investigators have now published EBC data on more than 100 healthy, full-term 4- and 5-month-olds. In addition, data collected from 14 premature

Table 2. Continued

| Task | Summary of test | Equipment required |
|------|-----------------|--------------------|
| Mobile/train conjugate reinforcement | Infants at 3 and 6 months of age are conditioned to move an overhead crib mobile by kicking one of their feet (mobile conjugate reinforcement). At 9 and 12 months, infants are conditioned to activate a musical train and a bank of 10 lights with a lever press response. At each age, 15-min conditioning sessions are conducted in a series of home visits separated by 24 hr. After conditioning sessions, infants are tested after increasing delays (1, 7, or 14 days later) until they exhibit no retention for 2 successive weeks. Contour detection and closure detection: Using the mobile described above, the infant learns to kick to move the mobile. After two learning sessions on 2 consecutive days, one or more visual characteristics (contour and closure) of the mobile are altered for some infants and not for others. On the third test day, recognition and discrimination of the test mobile are assessed using kick rates in the presence of the training mobile (old) or a novel mobile (new) relative to a baseline acquired for that infant before learning the task. | Stimulus: treat or toy Mobile or musical train with lighted press response box Limitations: Test is labor intensive Significant respondent burden Infant motivational factors also impact on test. The task cannot be automated, so problems associated with tester–subject interaction must be addressed. |
| Delayed nonmatching to sample (DNMS) | A sample object is presented. A delay follows, and then the familiar object is presented alongside a novel object. The correct choice is to select the novel object. | Object (toy or treat) |
| Means–end problem solving | At 7–8 months, task involves placing a cloth in reach of child and placing toy at the far end of cloth. To retrieve the toy, infant pulls the cloth (one-step problem solving). At 9 months, infants watch while toy is placed on end of cloth and then hidden under a cover. Infant has to first pull cloth to retrieve cover and then remove cover to find toy (two intermediate steps). At 10 months, infants must remove barrier to grasp cloth, pull cloth to retrieve cover, and search under cover to find toy (three intermediate steps). For each task, infants receive several trials to solve problem. Score is based on criteria for evidence of intention to retrieve the hidden toy (Willatts and Forsyth 2000). | Cloth to lay on tabletop Toy Cover to hide toy |
| Event-related potentials (ERPs) | Evaluation of a synchronized portion of the QEEG, time-locked to the onset of some event in the infant’s environment. | |
| Operant discrimination (object features and spatial mapping discrimination) | Visual/spatial displays are presented to the right and left of midline. Looking to a “correct” dimension (color, form, or spatial position) produces synchronous auditory reinforcement. Measures retention of correct dimension. | Displays, e.g., red circle, green square Auditory reinforcement: music Limitations: Tasks are not standardized for use to detect deficits in brain development or functioning. |
| Testing scales | Bayley Scales of Infant Development II: Individually administered instrument composed of two main subscales: mental scale, 118 items that assess mental ability (memory, habituation, problem solving, ability to vocalize, language and social skills); motor scale, 111 items that assess motor ability (rolling, crawling and creeping, sitting, standing, walking, running, jumping). All items arranged in order of developmental difficulty. Specification provided for specific sets of items to administer to a child depending on chronological age (Bayley 1993). Early Childhood Longitudinal Study reduced-item Bayley (ECLS-B): A reduced-item set developed that can be administered in less time and produce reliable, valid scores equivalent to the full set (West and Andreassen 2002). Items have been selected for their operational ease and psychometric properties. Multiple items can be scored from one administration, and, in the motor specialty, several items can be scored from observation. | 9-month-olds, approximately 25 min to administer. |
infants (28–31 weeks) using simple delay EBC have been submitted for publication (Herbert et al. In press).

**Animal model.** A rodent model for studying development of EBC is well established (Woodruff-Pak and Steinmetz 2000b). The emergence of EBC occurs gradually between 17 and 24 days of age in the rat. Disruption of cerebellar development by administering an antiproliferative agent, neonatal alcohol exposure, or early cerebellar or hippocampal aspirations interferes with development of normal EBC (Ivkovich and Stanton 2001; Stanton 2000; Stanton and Goodlett 1998).

Classical EBC represents a promising test of cognitive function with a well-studied homologous laboratory animal counterpart. Additional data are needed on population norms for infants and on the predictive validity or correlation of EBC deviations from established norms in infancy with later childhood and adult cognitive function assessments. Approaches to increasing subject retention rates between conditioning sessions and refinement of procedures to achieve higher success rates on criterion trials in each conditioning session will further strengthen this method.

**Visual Habituation/Novelty Preference Tasks and Visual Recognition Memory Tasks**

Tasks based on habituation/novelty and visual recognition memory (also called paired comparison) paradigms have been used widely to assess information processing and attention in infants and monkeys (Sirois and Mareschal 2002). Habituation occurs when attention decreases to repeated presentation of the same stimulus; novelty preference occurs when attention increases at the later presentation of a new stimulus. Infants and animals have a preference for novelty. Habituation and novelty preference are interpreted as reflecting the subject’s processing of stimulus information (Colombo 1993). Although the habituation/novelty paradigm focuses on the developmental course and speed with which attention wanes to a repeated stimulus, the visual recognition memory paradigm is concerned chiefly with visual recognition memory as reflected in differential responsiveness to familiar and novel stimuli. Such responsiveness is assessed after an initial exposure to the familiar stimulus, which is considerably briefer than that afforded in the habituation paradigm (Rose et al. 2001b).

**Paired-comparison task (look duration, shift rate, novelty score)—infant model.** In this task, the infant is presented with a target for a period of familiarization. When the familiar target is paired with a novel one, infants typically spend more time looking at the novel target, implying recognition memory. The examiner records the number of looks and looking time (Rose et al. 2001a).

**Habituation assessment—infant model.** In habituation studies, each trial is either fixed by the experimenter or determined by how long the infant keeps looking at a stimulus. The length of the intertrial interval may also be varied. Which aspect to use as a predictor of risk has been the focus of considerable debate (Colombo 1993; Fagen and Ohr 2001). A large body of evidence indicates that look duration is related to performance, such that infants with shorter looks process information faster and more efficiently than do infants with longer looks (Colombo 1993; Rose et al 2001a). In addition, short lookers tend to process global properties before local properties, much like adults do, whereas long lookers tend to focus initially on local aspects of the stimuli. Of course, there is no way to know whether equal look durations reflect equivalent depths of concentration, what is being encoded, or how rapidly it is being encoded (Rovee-Collier and Barr 2002).

The infant-control procedure represents an important evolution in visual habituation procedures (Lavoie and Desrochers 2002). In this procedure, a trial begins when the infant looks at the stimulus and ends when the infant looks away. In a study of the short-term reliability of this test, a number of habituation measures and reaction to novelty response were shown to be a reliable and valid construct.

**Visual recognition memory assessment—infant model.** There is substantial evidence that poorer performance on tests of visual recognition memory and slower habituation are associated with "risk" for cognitive delay. Among the groups studied are infants with Down syndrome and those with prenatal exposure to chemical teratogens, malnourishment, and prematurity (Rose and Orlian 2001). For example, in a recent longitudinal study of full-term and preterm (birth weight < 1,750 g) infants seen at 5, 7, and 12 months, full-term infants had shorter look durations, faster shift rates, less off-task behavior, and higher novelty scores than did preterms (Rose et al. 2001a).

Overall, mean predictive correlations are comparable for both habituation and visual recognition memory and tend to be approximately $r = 0.45$ (Rose and Orlian 2001). A prospective longitudinal study ($n = 109$) followed high-risk preterms and a socioeconomically matched group of full-terms annually through 6 years of age (Rose et al. 1992) and at age 11 (Rose and Feldman 1995). Visual recognition memory at 7 months and a 1-year cross-modal transfer (test of infant feeling object without seeing and then identifying it visually) each predicted Bayley scores at 2 years and IQ at 3, 4, 5, 6, and 11 years. Correlations of infancy scores with the various outcomes were similar for both groups and ranged from 0.37 to 0.65. Visual recognition memory and cross-modal transfer also correlated with speed of information processing, memory, and verbal and spatial abilities at 11 years of age (Rose et al. 1997).

**The Fagan Test of Infant Intelligence (FTII)—infant model.** The FTII is a standardized paired-comparison test developed in the 1980s for the early assessment of infant intelligence using the fixation preference principle (Fagan 1990a, 1990b; Fagan and Singer 1983). It has since been used to detect delayed mental development in infants subsequent to environmental exposure to neurotoxic chemicals (Darvill et al. 2000; Jacobson et al. 1985, 1996; Simmer 2000; Winneke et al. 1998). The test is constructed for use at four gestational ages, 67, 69, 70, and 92 weeks, corresponding to 27, 29, 39, and 52 weeks postnatal age. Reviews of the predictive validity of the FTII report correlations with later tests of intelligence at 36 months of age ranging from 0.31 to 0.61 (Fagan 1990a, 1990b; Fagan and Dettmer 1992). The instrument also correctly predicted more than 80% of infants who were later identified as mildly to severely retarded. FTII test results in the first year of life predict intellectual performance (Stanford-Binet Intelligence Scale IV; Thorndike et al. 1986) at 8 years of age (Smith et al. 2002). However, there are questions regarding the strength of predictive validity of the FTII in nonrisk samples and variability in correlations depending upon the infant’s age at testing (Andersson 1996).

Andersson (1996) found low predictive correlations (0.21) in a longitudinal study on a random sample of 100 boys and 96 girls assessed on the Fagan test at 7 and 9 months and then again at 5 years. Furthermore, retest reliabilities at 2-week intervals for two observers in a small nonrisk sample of children at 7 months of age were found to be zero or even slightly negative (Winneke et al. 1998). In addition, recent research has questioned whether recognition memory is what is being measured in tests of this type (Colombo 1993). Other cognitive factors that could affect the FTII and related tests include sensory or perceptual visual discrimination, or speed of visual processing. Premature infants do less well at 6 months and 12 months of age than do full-term infants (Rose 1983).

**Disengagement fixation task—infant model.** This task was designed to study whether individual and developmental differences in look duration are linked to development of neural attention systems that control the ability to disengage visual fixation (Frick et al. 1999). Look duration has been correlated with disengagement latency: longer-looking infants are slower than shorter-looking infants to shift fixation to a peripheral target on competition trials, but not on noncompetition trials. This task has been used only in a research setting examining the development of the
neural attention systems that control the ability to inhibit visual attention.  

**Span task—infant model.** The span task, based on visual recognition memory and paired comparisons, is designed to assess the amount of information infants can hold in short-term memory. Novelty scores provide a measure of performance on each task and an overall index of capacity (Rose et al. 2001b). Thus far, only one human study and no animal studies have used this task.

**Visual habituation/novelty preference tasks and visual recognition memory tasks—animal models.** In animals, the closest parallel tasks have been studied in monkeys. In the visual recognition memory test, adapted from human infant tasks described above, novel visual stimuli are paired with familiar stimuli and looking time for both is recorded. There are striking similarities between macaque monkeys and human infants in the development of visual recognition memory and other adaptations of paired-comparison tasks and in the effects of risks on cognition (Burbacher and Grant 2000). Monkey infants, like human infants, show deficits associated with severe birth trauma, exposure to teratogens, and low birth weight (Gunderson et al. 1987). Deficits in visual recognition memory have been documented, including exposure to methyl mercury (Gunderson et al. 1986), ethanol (Gunderson et al. 1987), and methanol (Burbacher and Grant 2000).

This task cannot be applied directly to rodents because the primary sensory modality is visual. Rat visual systems are relatively weak, and their “direction of gaze” is represented better by input from the auditory, tactile, or olfactory modalities (Bushnell 1998). Some have compared the novelty object proximity tasks in rats with the human infant paired-comparison tasks (Anderson 2000). This task measures the tendency of rats to explore an unfamiliar object placed within an open field. The limitations of applying tasks such as the novelty object proximity tasks and observational methods to studies of head gaze novelty preference in human infants and monkeys are reviewed by Bushnell (1998).

The A-not-B Task and the Delay-Tolerance A-not-B Tasks

Piaget’s A-not-B task is widely used to study infant cognitive development (Diamond 2001a). Under the name “delayed response,” the almost identical task is used in rhesus monkeys to study the functions of the dorsolateral prefrontal cortex (Goldman-Rakic 1987). Subjects must “hold in mind” for a few seconds where a treat (or toy) is hidden and, over trials, must update their mental record to record where the treat was hidden last. Subjects are rewarded for reaching correctly, hence reinforcing the response. This task requires an aspect of working memory (holding the information in mind) plus inhibition of the natural tendency to repeat a positively reinforced response on reverse trials.

**Infant model.** By roughly 7.5–8 months of age, human infants correctly reach the first hiding location with delays as long as 2–3 sec (Diamond 2001a, 2001b). When the reward is hidden at location B, infants make a mistake (called the A-not-B error) by going back again to the A hiding place. Between 7.5 and 12 months, infants show increasing improvements in their performance of the delayed-response A-not-B task. For example, each month they can withstand delays approximately 2 sec longer. By 12 months of age, delays of 10 sec or longer are needed to see the A-not-B error (Diamond 2001c).

Various adaptations of the procedures have been studied. In a longitudinal study on 13 infants, Bell and Fox (1992) rated infant’s performance on an ordinal scale. Infants proficient at reversal trials on a given day received a score corresponding to that level of delay. Investigators have also developed a looking version of the task in which the eye gaze, not the reach, is the criterion evaluated. No differences in the performance of more than 100 infants on the delay-tolerance A-not-B tasks with an eye-gaze response, compared with the reaching response, have been documented (Bell and Adams 1999).

Interobserver agreement ratings on the A-not-B task are reported in the range of 85–95%, with higher ratings where videotape is used (Bell and Adams 1999). Differences in task performance have been reported between normal control infants and infants with Down syndrome, autistic children, and cocaine-exposed infants (Noland 2001). There have been no demonstrations of predictive validity of the A-not-B task as a measure of individual difference (Noland 2001), although infants with phenylketonuria have been followed for 4 years with continued impaired performance on tests of frontal lobe functioning (Diamond 2001b).

Although there is a wealth of study and debate on establishing the cause of the response preservation seen in 8- to 12-month-olds in these tasks (Ahmed and Ruffman 1998; Carey and Xu 2001; Diamond 2001a, 2001b; Diedrich et al. 2001), a standardized procedure for the tasks has not been developed (Diamond 2001c; Noland 2001).

**Animal model.** Infant rhesus monkeys improve on these same tasks (more quickly reaching the hiding locations, withstanding longer delays) during the same equivalent age period—1.5–4 months (Diamond 1991). In monkeys, an adaptation of this task, the object concept test, has been used to study in utero exposure to methyl mercury, lead, and methanol (Burbacher and Grant 2000).

The Transparent Barrier Detour Task (Object Retrieval Task)—Infant and Animal Models

Like the delay-tolerance A-not-B task, this task has been used in human infants and in rhesus monkeys to study working memory and functions of the dorsolateral prefrontal cortex. At 6–8 months in human infants and the equivalent age in rhesus monkeys, subjects reach for a toy or treat in a clear box only at the side through which they are looking. As they get older, subjects can look through the opening, sit up, and reach in while looking through the closed side. Infants 11 or 12 months old and monkeys 4 months old do not need to look along the line of reach. Infants and monkeys progress through a well-demarcated series of five stages in performance of this task (Diamond 1991).

There are wide individual differences in the rate at which infants and monkeys move through the tasks to retrieve the object. However, the age at which a given subject achieves “phase 1B” on the object retrieval task is remarkably close to the age at which that same subject can first uncover a hidden object in the delayed-response A-not-B task. The object retrieval tasks and comparisons with performance of the delayed-response A-not-B task have been mainly studied in relation to development and function of the prefrontal cortex. The limitations described for the delayed-response A-not-B task also apply to the transparent barrier detour task.

**Mobile/Train Conjugate Reinforcement Tasks**

The mobile/train conjugate reinforcement tasks are based on operant conditioning and the rationale that infants who lack a verbal response can perform a motoric response (foot kick, lever press) to indicate whether they recognize a stimulus or reinforcement/ reward (Rovee-Collier and Barr 2002). The tasks involve acquisition of information regarding the relationship between behavior (kicking a foot or pushing a lever) and a reinforcement or reward (mobile or train moves). These tasks provide a direct means of assessing long-term memory (Fagen and Ohr 2001) because the extent to which the infant retains the learned action can be measured.

**Infant model.** Infants at 2–3 months and 6 months of age are conditioned to move an overhead crib mobile by kicking one of their feet, which is attached to the mobile by a ribbon. Foot kicks move the mobile in a graded manner that is commensurate with their rate and vigor, providing conjugate reinforcement. At 9 and 12 months, infants are conditioned to activate a musical train and a bank of 10 lights with a lever press response. At each age, 15-min conditioning sessions are conducted in...
Considerable reliability, validity, and population. The apparatus used to test monkeys (Paule 2000) the OTB and has been studied in children 6.5 and attention (Diamond et al. 1999; Paule et al. 1992, 1993). Likewise, infant monkeys do not reliably reach criterion on DNMS at 10-sec delays until 4 months of age. Diamond et al. (1999) postulated that infants failed on the DNMS not because of lack of memory requirements, but because infants did not understand the relationship between stimulus and reward or because spatial separations between response and reward or between stimulus and response make the task more difficult.

Diamond et al. (1999) have designed a DNMS task for infants in which the infants do not displace stimuli to receive rewards, but the objects used as stimuli themselves are the reward. The protocol is the same as for the DNMS except the rewards are attached to the base of the stimuli. With this modification, 70% of 9-month-olds succeeded in the DNMS with a 5-sec delay. When verbal rewards (experimenter cheered and applauded when the infant reached correctly) were provided, 80% of infants passed the DNMS with a 5-sec delay (Diamond et al. 1999). Diamond and colleagues' modification of the standard DNMS has not been well studied as an assessment tool in infants, although there is a growing body of data on the standard DNMS in older children (Chelonis et al. 2000).

**Means–End Problem-Solving Task**

Means–end problem solving involves the deliberate and planned execution of a sequence of steps to achieve a goal. Means–end behavior develops after 6 months of age and involves the acquisition of knowledge of appropriate means–end relations and abilities such as planning, sequencing actions, and maintenance of attention to a goal (Willatts and Forsyth 2000). There is evidence that development of means–end problem solving is related to development of the prefrontal cortex (Diamond et al. 1997).

**Infant model.** Infants between 7 and 8 months of age can solve simple problems involving the completion of one intermediate step—for example, pulling a cloth to retrieve a toy sitting on top of it. By 9 months, infants begin to solve more complex problems requiring completion of two intermediate steps to achieve a goal. Infants first watch as a toy is placed at the end of a cloth and then hidden by a cover. To solve the problem, an infant must first pull the cloth to retrieve the cover and next remove the cover to find the toy. At 10 months, infants can solve more complex problems involving three intermediate steps: removing a barrier to grasp a cloth, pulling the cloth to retrieve a cover, and searching under the cover to find a toy (Willatts 1999). Means–end problem-solving tasks are structured so that the infant’s sequence of behavior is scored according to specified criteria for evidence of intention to retrieve the hidden toy, with higher scores indicating more mature problem solving (Willatts and Forsyth 2000).

Two-step problem-solving scores at 9 months of age correlate positively with IQ (0.64, p < 0.01) and vocabulary scores (0.42, p < 0.01) at 3 years (Slater 1995; Willatts 1997). In a randomized trial of the role of long-chain polyunsaturated fatty acids in infant cognitive development, higher problem-solving scores were observed on the 9-month two-step problem-solving task in the supplemented infants who at 3 months had demonstrated poorer attention control and had a lower birth weight. At 10 months, all children in the supplemented group displayed higher problem-solving scores on the three-step task (Willatts and Forsyth 2000).

**Animal model.** The incremental repeated acquisition (IRA) task, part of the OTB, might be considered a parallel test in monkeys and rodents. The animal is required to learn a sequence of lever presses to receive a reinforcer. First, in IRA1, the subject is required to learn the correct response to one of three levers. Next, in IRA2, the subject is required to learn a response on a different lever than for IRA1, and then a two-lever sequence. The tasks are incremented up to a six-lever sequence or until the allotted task time has elapsed (Mayorga et al. 2000a).

**Event-Related Potentials**

Quantitative electroencephalographic (QEEG) measures have been used in clinical settings to diagnose neuropathology and, in infants, to evaluate gestational age and maturational levels of newborns. The use of electroencephalographic (EEG) recordings in conjunction with other task measures has become a common practice in studying psychophysiological processes. The use of EEG measures in conjunction with A-not-B tasks is reviewed by Marshall and Fox (2001).

An event-related potential (ERP) is a synchronized portion of the ongoing EEG pattern. The ERP is distinguished from the more traditional baseline EEG measure in that the evoked potential is a portion of the ongoing EEG activity that is time-locked to the onset of some event in the infant’s environment (Molfese and Molfese 2001). The ERP reflects both general and specific aspects of the evoking stimulus and the person’s perceptions and decisions regarding it (cognition) as reflected by changes in the amplitude or height of the wave at different points in its time course. ERPs are recognized as providing information concerning between-hemisphere differences as well as within-hemisphere differences in the brain’s electrical activity under specific stimulus conditions.
Infant model. ERPs have been paired with both vision and auditory assessments in infants and correlated with later intelligence measures. Studies in the 1960s through the 1980s using ERPs had mixed results (Molfese and Molfese 2001). Recent studies on small samples using newer technology and improved study design suggest that ERPs have value as predictors of later functioning. Studies reviewed by Molfese and Molfese (2001) showed measures obtained in later infancy and early childhood successfully predicted language and cognitive skills in older children. Nelson et al. (2000) used ERPs paired with auditory stimuli to test auditory recognition memory in normal newborn infants and the infants of diabetic mothers. Neonatal ERP’s elicited by the maternal voice were compared with those elicited by a stranger’s voice. Results were compared with Bayley scores at 1 year of age. The presence of a specific neonatal ERP pattern (greater positive slow wave area in response to stranger’s voice) indicated better 1-year cognitive development. In an earlier study (Nelson and Bloom 1997), ERPs were used for shape recognition at 4 months in high-risk preterm infants and healthy full-term infants. ERPs were recorded while infants were familiarized with one stimulus (a red cross, 15 trials) and a novel stimulus (red corkscrew). Atypical patterns were found in the high-risk infants.

Animal model. ERPs can be recorded in monkeys (Lilienthal and Winneke 1996; Lilienthal et al. 1994) and rodents (Winneke 1992) and are being used in a parallel paired-comparison task in both monkeys and human infants in the University of Michigan longitudinal study of iron deficiency study (Lozoff B. Personal communication). Rhesus monkeys pre- and postnatally exposed to lead had consistent prolongations of latencies in the brainstem auditory evoked potentials (Lilienthal and Winneke 1996) and visually evoked potentials (Lilienthal et al. 1988).

Operant Discrimination Learning (Object Features and Spatial Mapping Discrimination Tasks)

Infant model. Colombo (2001) trained 3-, 6-, and 9-month-olds to an association between an auditory reinforcement and attention to visual/spatial displays. Colombo (2001) reviewed similar studies by Harman et al. (1994) and work by Catherwood et al. (1996) on determining the time course of the processing of visual features and their joining compounds in 5- to 6-month-olds. It is important to note that tasks such as these are currently used to examine how the infant brain develops and functions.

Animal model. Discrimination tasks in nonhuman primates are homologous to this task. The spontaneous alteration task in rats maybe a parallel model for this task. Rats exposed to PCBs prenatally showed altered performance on retention of visual discrimination tasks (Lilienthal and Winneke 1991).

Bayley Scales of Infant Development II

Infant model. The Bayley scales are considered the gold standard for assessing the current developmental functioning of infants and children from 1 to 42 months old. The Bayley Scales of Infant Development II have two main subscales, or sets of items: the 178-item mental scale and the 111-item motor scale (Bayley 1993). The mental and motor scales are known to be well correlated, because several items are scored for both scales. A reduced set of Bayley items has been developed that can be administered in less time and produces reliable, valid scores equivalent to those of the full set (West and Andreassen 2002). The Early Childhood Longitudinal Study reduced-item Bayley (ECLS-B) for 9-month-olds takes approximately 25 min to administer. Items have been selected for their operational ease and psychometric properties. Multiple items can be scored from one administration, and, in the motor scale especially, several items can be scored from observation.

Animal model. Adaptations of many of the assessments included in the Bayley Scales for Infant Development II (Bayley 1993) have been used in assessment of nonhuman primate infants (Burkhauser and Grant 2000).

Discussion

Over the last decade, there have been tremendous advances in the understanding of the development of learning, memory, and attention in infants and in measures to assess these functions. This review identifies validated tests of normal cognitive function in human infants 12 months and younger that have a homologous or parallel test in laboratory animals. The tests vary along many dimensions of desirable properties, including the number of available validation data, speed of testing, breadth and/or specificity of test results, requirements for equipment and personnel to conduct the test, and extrapolation of results among species. As technology improves our ability to study infant brain function, continued advances are anticipated in the understanding of the integration of motor and mental development in infant cognitive function, and in identifying the factors that arrest normal development.

Tests are currently under study in human infants that seem appropriate for evaluation in animal models. For example, the visual expectation paradigm (VEXP) is based on the infant’s ability to learn a spatiotemporal pattern (Haith et al. 1993). VEXP measures of expectation (anticipations and reaction times) have a moderate amount of reliability and stability (Canfield et al. 1995). Correlations of –0.44 to –0.46 between reaction time in infants (3.5 and 8 months) and standardized IQ measures at 3 to 4 years of age have been reported (DiLella et al. 1990; Dougherty and Haith 1997).

Some neurophysiologic tests also have promise in future research of brain function and cognitive development, such as functional magnetic resonance imaging and positron emission tomography. At this time, practical limitations restrict use to older children, because the tests require an alert, cooperative child who can overcome the fear of a strange situation and hold his or her head still (Singer 2001). Because a gold standard does not exist for assessment of cognitive function in animals for comparison with human infants, a battery of tests may also be considered. For example, B. Lazof at the University of Michigan (personal communication) is leading a longitudinal study assessing cognitive and motor development in 150 human infants and monkeys with iron deficiency from 9 to 12 months of age. The battery of tests assessing cognitive function include the A-not-B task, a Fagan II novel preference test modified to include looking time, the Resnick ocular motor spatial task, the pair-comparison task with ERP recordings, and a spatial recognition task.

Although sensorimotor and language development in infancy obviously prevents the assessment of late-maturing higher order skills that might be particularly sensitive to neurotoxicant exposures (e.g., reading, complex problem solving, executive functions such as planning, organizing, and strategizing skills), the concurrent validity of habituation and classical conditioning tests has been established. Bellinger (2002) suggests interpreting the validated infant assessment tests in the same way that neonatologists interpret birth weight. Although not predictive of later weight, birth weight is highly informative as an index of a newborn’s general health status.

An important advantage of assessing cognitive outcomes in infancy is reducing the amount of time between gestational or early postnatal neurotoxicant exposure and outcome assessment. This has several applications. Early assessment increases the strength of, and reduces the bias in, the estimate of the neurotoxicant’s contribution to the results of the neurobehavioral assessment. By reducing the time available for other factors to influence outcome, early assessment allows observation of the relatively direct effect of the exposure. Conversely, but of perhaps equal importance, results obtained in childhood are assessed longitudinally to help identify the impact on neurodevelopment of confounding factors (e.g., sociodemographic, education). This information can be valuable for developing intervention strategies, which in turn are often more effective when initiated earlier in development.
Comparative measures in laboratory ani-
ma ls are essential to the understanding of
the tox icology underlying effects measured in
human infants. Assessment of the actual risk to
developing humans relies heavily on extrapola-
tion of data from animal studies. When com-
parable methods are used in laboratory studies
and in evaluation of human infants, more con-
fidence can be placed on predictions of levels
of exposure that will adversely affect humans.
Ethical and economic considerations support
the choice of rodent over nonhuman primate
studies. Homologous measures, in which the
identical methodology is employed, have some
advantages over parallel measures, which rely
on different techniques to evaluate what are
believed to be the same processes in different
species. We found several homologous tasks for
humans and nonhuman primates, some suit-
able for the study of infants. We identified
only one homologous task, classic EBC, that
can be used in humans, nonhuman primates,
and rodents. It is not sufficiently developed
at this time for use in large-scale studies.
However, we identified several parallel me-
asures that are suitable for evaluation of human
infants and application in rodent toxicology
studies designed to clarify and extend the
findings of studies in humans.

REFERENCES

Adams J, Barone S Jr, LaMantha A, Philen R, Rice DC, Spear L, et al. 2000. Workshop to identify critical windows of expo-
sure for children's health: neurobehavioral work group summary.
Environ Health Perspect 108(suppl 1):335–344.
Ahmed A, Ruffman T. 1998. Why do infants make and B errors in
a search task, yet show memory for the location of hid-
den objects in a nontarget task? Dev Psychol 34:441–453.
Andersson BK. 2000. The factor in non-human animals. Novartis
Found Symp 232:79–90.
Andersson HW. 1996. The Fagan Test of Infant Intelligence:
predictive validity in a random sample. Psychol Rep 78:1059–1062.
Bayley N. 1993. Bayley Scales of Infant Development Manual,
2nd ed. San Antonio, TX:Psychological Corporation, Harcourt Brace and Company.
Bell MA, Adamson SE. 1998. Comparable performance on looking and
reaching versions of the A-not-B task at 8 months of
age. Infant Behav Dev 21:221–235.
Bell MA, Fox NA. 1992. The relations between frontal brain
electrical activity and cognitive development during infancy.
Child Dev 63:1142–1163.
Bellinger DC. 2002. Perspectives on incorporating human neu-
robehavioral end points in risk assessments. Risk Anal 22:467–498.
Burghuber TM, Grant KS. 2000. Methods for studying nonhuman
primates in neurobehavioral toxicology and teratology.
Neurotoxicol Teratol 22:475–486.
Bushnell PJ. 1998. Behavioral approaches to the assessment of
attention in animals. Psychopharmacologia (Berl) 138:231–259.
Bushnell PJ, Rice DC. 1999. Behavioral assessments of learning and
attention in rats exposed perinatally to 3,3',4,4'-pento-
tachlorobiphenyl (PCB 126). Neurotoxicol Teratol 21:381–392.
Campbell BA, Campbell EH. 1982. Retention and extinction
of learned fear in infant and adult rats. J Comp Physiol Psychol 55:1–8.
Campbell BA, Coulter X. 1976. Neural and psychological
processes underlying the development of learning and
memory. In: Habituation (Tighe J, Leaton RN, eds). Hillsdale, NJ:Lawrence Erlbaum, 129–157.
Canfield RJ, Wilkin JL, Schmerl L, Smith EG. 1995. Age-related
change and stability of individual differences in infant sac-
cade reaction time. Infant Behav Dev 18:351–358.
Carey S, Xu F. 2001. Infants' knowledge of objects: beyond
object files and object tracking. Cognition 80:179–213.
Carlson K, Dawson VL. 1996. Assessing the primary
moms in infant encoding of compound visual
stimuli. Infant Behav Dev 19:1–11.
Chelions JI, Daniels-Shaw JL, Blake DJ, Paulie MG. 2000.
Developmental trajectories (Maccabe) of sample task
performance in children. Neurotoxicol Teratol 22:683–694.
Colombo J. 1993. Infant Cognition: Predicting Later Intellectual
Functioning. Newbury Park, CA:Sage.
———. 2001. The development of visual attention in infancy.
Annu Rev Psychol 52:327–367.
Cory-Slehta DA, Crofton KM, Foran JA, Ross JF, Shepts LP,
Weiss B, et al. 2001. Methods to identify and characterize
developmental models for human risk assess-
ment I: behavioral effects. Environ Health Perspect 109(suppl 1):79–91.
Davill T, Lenkey E, Reiman J, Stewart P, Pagano J. 2000.
Prenatal exposure to PCBs and infant performance on the Fagan Test of Infant Intelligence.
Neurotoxicol 21:1029–1038.
Diamond A. 1990. Rate of maturation of the hippocampus and
the developmental progression of children's performance on the delayed non-matching to sample and visual paired
comparison tasks. Ann NY Acad Sci 608:394–426.
———. 1991. Neuropsychological insights into meaning of
object concept development. In: The Epigenesis of Mind:
Essays on Biology and Cognition (Carey S, Gelman R, eds).
Hillsdale, NJ:Lawrence Erlbaum, 67–110.
———. 2001a. Prefrontal cortex development and develop-
cement of cognitive function. In: International Encyclopedia
of the Social and Behavioral Sciences (Smelser NJ, Baltes PB, eds).
Oxford, UK:Elsevier Science, 11769–11892.
———. 2001b. A model system for studying the role of dopamine
in the prefrontal cortex during early development in humans:
early and continuously treated phenothiazines. Handbook of
Developmental Cognitive Neuroscience (Nelson CA, Luciana M, eds). Cambridge, MA:MIT Press, 433–472.
———. 2001c. Looking closely at infants' performance and
experimental procedures in the A-not-B task. Behav Brain
Sci 24:38–41.
Diamond A, Churchland A, Cruess L, Kirkham NZ. 1999. Early
developments in the ability to understand the relation
between stimulus and reward. Dev Psychol 25:1607–1617.
Diamond A, Prevor MB, Callender G, Druin DP. 1997. Prefrontal
Cortex Cognitive Deficits in Children Treated Early and Continuously for PKU. Waltham, MA:Center for
Developmental Cognitive Neuroscience.
Diedrich FC, Highman TD, Spahr KA, Theilen E, Smith LB. 2001.
The role of target distinctiveness in infantive preservative
rearing. J Exp Child Psychol 78:263–290.
Dill JF, Thompson IE, Kohler K, Turfield MJ, Prever FL, Jaffe HM, Matthiessen JJ, Baltes PB, et al. 1999. A technical report. J Appl
Dev Psychol 13:153–157.
Dougherty TM, Hartmann M. 1997. Infant expectations and reac-
tion times as a function of speed of processing and
IQ. Dev Psychol 33:146–155.
Fagan JF III. 1990a. The paired-comparison paradigm and infant
intelligence. In: The Development and Neural Bases of
Higher Cognitive Function (Diamond A, ed). New York/New
York Academy of Sciences Press, 337–364.
Fagan JF, Detterman D. 1992. The Fagan Test of Infant Intelligence:
a technical report. J Appl Dev Psychol 13:153–157.
Fagan JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JW, Ohr PS. 1990. Individual differences in infant condi-
tional memory and conditioning. In: Memory Differences in
Infancy. Presented at the 3rd Annual Meeting of the Society for
Neurosciences, November 1989, Washington, DC.
Fagen JF, Detterman D. 1992. The Fagan Test of Infant Intelligence:
A model of infant memory and intelligence. J Exp Child Psychol 78:263–290.
Fagan JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Detterman D. 1992. The Fagan Test of Infant Intelligence:
a technical report. J Appl Dev Psychol 13:153–157.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
Fagen JF, Singer LT. 1983. Infant recognition memory as a meas-
ure of intelligence. In: Advances in Infancy Research, Vol 2
(Lipsett LP, ed). Norwood, NJ: Ablex, 31–78.
and language processes in infancy. In: Biobehavioral Assessment of the Infant (Singer LT, Zeskind PS, eds). New York:Guilford, 323–340.

Nelson CA, Bloom FF. 1997. Child development and neuroscience. Child Dev 68:970–987.

Nelson CA, Luciana M, eds. 2001. Handbook of Developmental Cognitive Neuroscience. Cambridge, MA:MIT Press.

Nelson CA, Wewerska S, Thomas KM, deRegnier RA, Tribbey-Walbridge S, Georgieff M. 2000. Neurocognitive sequelae of infants of diabetic mothers. Behav Neurosci 114:950–956.

Noland JS. 2001. The A-not-B task. In: Biobehavioral Assessment of the Infant (Singer LT, Zeskind PS, eds). New York:Guilford, 312–322.

Overman WH. 1990. Performance on traditional matching to sample, non-matching to sample, and object discrimination tasks by 12- to 32-month-old children. A developmental progression. Ann NY Acad Sci 608:365–385.

Overman WH, Bachevalier J. 2001. Inferences about the functional development of neural systems in children via the application of animal tests of cognition. In: Handbook of Developmental Cognitive Neuroscience (Nelson CA, Luciana M, eds). Cambridge MA:MIT Press, 109–124.

Overman W, Bachevalier J, Miller M, Moore K. 1996a. Children’s performance on “animal tests” of oddity: implications for cognitive processes required for tests of oddity and delayed nonmatch to sample. J Exp Child Psychol 62:223–242.

———. 1996b. Cognitive gender differences in very young children parallel biologically based cognitive gender differences in monkeys. Behav Neurosci 110:673–684.

Overman WH, Bachevalier J, Sewell F, Drew J. 1993. A comparison of children’s performance on two recognition memory tasks: delayed nonmatch-to-sample versus visual paired-comparison. Dev Psychobiol 26:345–357.

Overman W, Bachevalier J, Turner M, Peuster A. 1982. Object recognition versus object discrimination: comparison between human infants and infant monkeys. Behav Neurosci 106:15–29.

Paule MG. 2000. Validation of a behavioral test battery for monkeys. In: Methods of Behavioral Analysis in Neuroscience (Buccafusco JJ, ed). Boca Raton, FL:CRC Press, 281–294.

Paule MG. 1999a. Operant test battery performance in children: correlation with IQ. Neurotoxicol Teratol 21:223–230.

Paule MG, Popke EJ, Pearson E, Hammond T. 1999b. Development of a nonhuman primate model for studying the consequences of long-term neuroprotectant administration on complex brain functions in developing animals. Ann NY Acad Sci 890:470.

Dick B, Barone S Jr. 2000. Critical periods of vulnerability for the developing nervous system: evidence from humans and animal models. Environ Health Perspect 108:suppl 3:511–533.

Rose SA. 1983. Differential rates of visual information processing in full-term and preterm infants. Child Dev 54:1198–1198.

Rose SA, Feldman JF. 1995. Prediction of IQ and specific cognitive abilities at 11 years from infancy measures. Dev Psychol 31:685–696.

Rose SA, Feldman JF, Futterweit LR, Jankowski JJ. 1997. Continuity in visual recognition memory: infancy to 11 years. Intelligence 24:381–392.

———. 2001a. Visual short-term memory in the first year of life: capacity and recency effects. Dev Psychol 37:329–549.

Rose SA, Feldman JF, Wallace IF. 1992. Infant information processing in relation to six-year cognitive outcomes. Child Dev 63:1126–1141.

———. 2001b. Visual short-term memory in the first year of life: capacity and recency effects. Dev Psychol 37:329–549.

Rose SA, Orlian EK. 2001. Visual information processing. In: Biobehavioral Assessment of the Infant (Singer LT, Zeskind PS, eds). New York:Guilford, 274–292.

Rovee-Collier C, Barr R. 2002. Infant cognition. In: Stevens’ Handbook of Experimental Psychology (Pashler H, Wixted J, eds). New York:Guilford, 274–292.

Rose SA, Feldman JF, Wallace IF. 1992. Infant information processing in relation to six-year cognitive outcomes. Child Dev 63:1126–1141.

Singer LT. 2001. General issues in infant assessment and development. In: Biobehavioral Assessment of the Infant (Singer LT, Zeskind PS, eds). New York:Guilford, 274–292.

Sirois S, Mareshcal D. 2002. Models of habitation in infancy. Trends Cogn Sci 6:293–298.

Slater A. 1995. Individual differences in infancy and later IQ. J Child Psychol Psychiatry 36:69–112.

Sliker W Jr, Beck BD, Cory-Slechta DA, Paule MG, Anger WK, Bellinger D. 2000. Cognitive tests: interpretation for neurotoxicity? Toxicol Sci 58:222–234.

Smith L, Fagan FF, Ulvand SE. 2002. The relation of recognition memory in infancy and parental socioeconomic status to later intellectual competence. Intelligence 30:247–259.

Stanton ME. 2000. Multiple memory systems, development and conditioning. Behav Brain Res 110:25–37.

Stanton ME, Freeman JH Jr. 1994. Eyeblink conditioning in the infant rat: an animal model of learning in developmental neu-rotoxicology. Environ Health Perspect 102:suppl 2:131–139.

Stanton ME, Goodlett CR. 1998. Neonatal ethanol exposure impairs eyeblink conditioning in weaning rats. Alcohol Clin Exp Res 22:270–275.

Stanton ME, Spear LP. 1990. Workshop on the qualitative and quantitative comparability of human and animal developmental neurotoxicity, Work Group I report: comparability of measures of developmental neurotoxicity in humans and laboratory animals. Neurotoxicol Teratol 12:261–267.

Thordnile RK, Hagen EF, Sattler JM. 1986. Stanford-Binet Intelligence Scale. Technical Manual. 4th ed. Chicago, IL:Riverside.

U.S. EPA. 1998. Guidelines for neurotoxicity risk assessment. Fed Reg 63(30):26926–26954.

Walters P, Forsyth JS. 2000. The role of long-chain polyunsaturated fatty acids in infant cognitive development. Prostaglandins Leukot Essent Fatty Acids 63:95–100.

Willatts P. 1997. Beyond the “couch potato” infant: how infants use their knowledge to regulate action, solve problems, and achieve goals. In: Infant Development: Recent Advances (Brenner J, Slater A, Butterworth G, eds). Hove, East Sussex, UK:Psychology Press, 109–135.

———. 1999. Development of means-end behavior in young infants: pulling a support to retrieve a distant object. Dev Psychol 35:651–667.

Willatts P, Forsyth JS. 2000. The role of long-chain polyunsaturated fatty acids in infant cognitive development. Prostaglandins Leukot Essent Fatty Acids 63:95–100.

Winneke G. 1992. Cross species extrapolation in neurotoxicology: neurophysiological and neurobehavioral aspects. Neurotoxicology 13(1):15–25.

Wonneke G, Buchalski A, Heinzow B, Kramer U, Schmidt E, Waldowial J, et al. 1998. Developmental neurotoxicity of polychlorinated biphenyls (PCBs): cognitive and psychomotor functions in 7-month-old children. Toxicol Lett 102:423–428.

Woodruff-Pak D, Steinmetz J, eds. 2000a. Eyeblink Classical Conditioning, Vol 2: Applications in Humans. New York:Kluwer Academic.