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VALIDITY AND RELIABILITY OF QUANTITATIVE ELECTROENCEPHALOGRAPHY (qEEG)

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ABSTRACT

Reliability and validity are statistical concepts that are reviewed and then applied to the field of quantitative electroencephalography or qEEG. The review of the scientific literature demonstrated high levels of split-half and test re-test reliability of qEEG and convincing content and predictive validity as well as other forms of validity. qEEG is distinguished from non-quantitative EEG (“Eye Ball” examination of EEG traces) with the latter showing low reliability (e.g., 0.2 to 0.29) and poor inter-rater agreement for non-epilepsy evaluation. In contrast, qEEG is greater than 0.9 reliable with as little as 40 second epochs and remains stable with high test re-test reliability over many days and weeks. Predictive validity of qEEG is established by significant and replicable correlations with clinical measures and accurate predictions of outcome and performance on neuropsychological tests. In contrast, non-qEEG or “Eye Ball” visual examination of the EEG traces in cases of non-epilepsy has essentially zero predictive validity. Content validity of qEEG is established by correlations with independent measures such as the MRI, PET and SPECT, the Glasgow Coma Score, neuropsychological tests, etc. where the scientific literature again demonstrates significant correlations between qEEG and independent measures known to be related to various clinical disorders. In contrast, non-qEEG or “Eye Ball” visual examination of the EEG traces in cases of non-epilepsy has essentially zero content validity. The ability to test and evaluate the concepts of reliability and validity are demonstrated by mathematical proof and simulation where one can demonstrate test re-test reliability for themselves as well as zero physiological validity of coherence and phase differences when using an average reference and Laplacian montage.

Key Terms: Quantitative EEG, Reliability, Validity

Quantitative electroencephalography (qEEG) is distinguished from visual examination of EEG traces, referred to as “non-quantitative EEG” by the fact that the latter is subjective and involves low sensitivity and low inter-rater reliability for non-epilepsy cases (Cooper et al, 1974; Woody, 1966; 1968; Piccinelli et al, 2005; Seshia et al, 2008; Benbadis et al, 2009; Malone et al, 2009). In contrast, the quantitative EEG (qEEG) involves the use of computers and power spectral analyses and is more objective with higher reliability and higher clinical sensitivity than is visual examination of the EEG traces for most psychiatric disorders and traumatic brain injury (Hughes and John, 1999). The American Academy of Neurology draws a distinction between digitization of EEG for the purposes of visual review versus quantitative EEG which is defined as: “The mathematical processing of digitally recorded EEG in order to highlight specific waveform components, transform the EEG into a format or domain that elucidates relevant information, or associate numerical results...” (Nuwer, 1997, p. 2). Thus, the definition of quantitative EEG is very broad and pertains to all spectral measures and numerical analyses including coherence, power, ratios, etc.

The low reliability of visual examination of EEG traces has been known for many years (Woody, 1968a; 1968b). As stated in a recent visual non-qEEG study by Malone et al (2009, pg. 2097):

“The interobserver agreement (Kappa) for doctors and other health care professionals was poor at 0.21 and 0.29, respectively. Agreement with the correct diagnosis was also poor at 0.09 for doctors and -0.02 for other healthcare professionals.”

Or in a study of non-qEEG visual examination of the EEG traces it was concluded by Benbadis et al (2009, pg. 843): “For physiologic nonepileptic episodes, the agreement was low (kappa = 0.09)”

A recent statement by the Canadian Society of Clinical Neurophysiology further emphasizes the low reliability of visual examination of EEG traces or non-qEEG in the year 2008 where they conclude:

“A high level of evidence does not exist for many aspects of testing for visual sensitivity. Evidenced-based studies are needed in several areas, including (i) reliability of LED-based stimulators, (ii) the most appropriate montages for displaying responses, (iii) testing during pregnancy, and (iv) the role of visual-sensitivity testing in the diagnosis of

neurological disorders affecting the elderly and very elderly.” (Seshia et al, 2008, pg. 133).

The improved sensitivity and reliability of qEEG was first recognized by Hans Berger in 1934 when he performed a qEEG analysis involving the power spectrum of the EEG with a mechanical analog computer and later by Kornmuller in 1937 and Grass and Gibbs (1938) (see Niedermeyer and Lopes Da Silva, 2005). qEEG in the year 2010 clearly surpasses conventional visual examination of EEG traces because qEEG has high temporal and spatial resolution in the millisecond time domain and approximately one centimeter in the spatial domain which gives qEEG the ability to measure network dynamics that are simply “invisible” to the naked eye. Over the last 40 years the accuracy, sensitive, reliability, validity and resolution of qEEG has steadily increased because of the efforts of hundreds of dedicated scientists and clinicians that have produced approximately 90,000 qEEG studies cited in the National Library of Medicine’s database . The estimate of 90,000 studies is from sampling of abstracts from the larger universe of 103,230 citations which includes both non-quantitative and quantitative EEG studies. The search term “EEG” is necessary because the National Library of Medicine searches article titles and rarely if ever is the term “qEEG” used in the title (e.g., this author has published over 150 peer reviewed articles on qEEG and has never used the term “qEEG or QEEG” in the title). Since approximately 1975 it is very difficult to publish a non-qEEG study in a peer reviewed journal because of the subjective nature of different visual readers agreeing or disagreeing in their opinions about the squiggles of the “EEG” with low “Inter-Rater Reliability” for non-epilepsy cases (Cooper et al, 1974; Woody, 1966; 1968; Piccinelli et al, 2005; Seshia et al, 2008; Benbadis et al, 2009; Malone et al, 2009). In this paper, I will not discuss the issue of qEEG in the detection of epilepsy. This topic is well covered by many studies (see Niedermeyer and Lopes Da Silva, 2005). Instead, this paper is focused on the non-epilepsy cases, the very cases that visual non-qEEG is weakest. It is useful to first re-visit the standard concepts of “Reliability” and “Validity” of quantitative EEG while keeping in mind the historical background of non-qEEG visual examination of EEG traces which is used in approximately 99% of the U.S. hospitals as the accepted standard of care in the year 2010 even though non-qEEG is insensitive and unreliable for the evaluation of the vast majority of psychiatric and psychological disorders and mild traumatic brain injury. Given this background, the purpose of this paper is to define the concepts of “Reliability” and “Validity” and evaluate these concepts as they apply

to the clinical application of qEEG. Such an endeavor requires some knowledge of the methods of measurement as well as about the basic neuroanatomy and neurophysiology functions of the brain.

It is not possible to cover all clinical disorders and therefore mild traumatic brain injury will be used as examples of qEEG validity and reliability. The same high levels of clinical validity and reliability (i.e., > 0.95) of qEEG have been published for a wide variety of psychiatric and psychological disorders to cite only a few, for example, attention deficit disorders (Mazaheri et al, 2010; van Dongen-Boomsma et al, 2010), ADHD (Gevensleben et al, 2009); Schizophrenia (Siegle et al, 2010; Begić et al, 2009); Depression (Pizzagalli et al, 2004); Obsessive compulsive disorders (Velikova et al, 2010); addiction disorders (Reid et al, 2003); anxiety disorders (Hannesdóttir et al, 2010) and many other disorders. The reader is encouraged to visit the National Library of Medicine database at:

<https://www.ncbi.nlm.nih.gov/sites/entrez?db=pubmed> and use the search terms “EEG and xx” where xx = a clinical disorder. Read the methods section to determine that a computer was used to analyze the EEG which satisfies the definition of quantitative electroencephalography (qEEG) and then read the hundreds of statistically significant qEEG studies for yourself. Because non-significant studies are typically not published it is no surprise that all of the clinical studies that this author read in the National Library of Medicine database were statistically valid and reliable. I was unable to find any clinical studies that stated that qEEG was not valid or not reliable. This is the same conclusion drawn by Hughes and John (1999).

Validity Defined

Validity is defined by the extent to which any measuring instrument measures what it is intended to measure. In other words, validity concerns the relationship between what is being measured and the nature and use to which the measurement is being applied. One evaluates a measuring instrument in relation to the purpose for which it is being used. There are three different types of validity: 1- Criterion-related validity also called “Predictive Validity”, 2- Content validity also called “face validity” and, 3- Construct validity. If a measurement is unreliable then it can not be valid, however, if a method is reliable it can also be invalid, i.e., consistently off the mark or consistently

wrong. Suffice it to say that clinical correlations are fundamental to the concept of validity and are dependent on our knowledge of basic neuroanatomy and neurophysiology. These concepts are also dependent on our methods of measurement and the confidence one has in the mathematical simulations when applied in the laboratory or clinical context. Today there are a wide number of fully tested mathematical and digital signal processing methods that can be rapidly evaluated using calibrated signals and a high speed computer to determine the mathematical validity of any method and I will not spend a lot of time on this topic except for a brief mention of a few methods that are not valid when applied to coherence and phase measures because of technical limitations, for example, the use of an average reference or the Laplacian surface transform and Independent Components Analysis (ICA) and the calculation of coherence and phase. It will be shown in a later section that the average reference and the Laplacian distort the natural physiological phase relationships in the EEG and any subsequent analyses of phase and coherence are invalidated when these remontaging or reconstruction methods are used (Rappelsberger, 1989; Nunez, 1981). The average reference and Laplacian and ICA methods are valid for absolute power measures but have limitations for phase measures which is a good example of why validity is defined as the extent to which a measuring instrument measures what it is intended to measure.

Leaving the mathematical and simulation methods aside for the moment, the most critical factor in determining the clinical validity of qEEG is knowledge about the neuroanatomy and neurophysiology and functional brain systems because without this knowledge then it is not possible to even know if a given measurement is clinically valid in the first place. For example, neurological evaluation of space occupying lesions has been correlated with the locations and frequency changes that have been observed in the EEG traces and in qEEG analyses, e.g., lesions of the visual cortex resulted in distortions of the EEG generated from the occipital scalp locations or lesions of the frontal lobe resulted in distortions of the EEG traces arising in frontal regions, etc. However, early neurological and neuropsychological studies have shown that function was not located in any one part of the brain (Luria, 1973). Instead the brain is made up of complex and interconnected groupings of neurons that constitute “functional systems”, like the “digestive system” or the “respiratory system” in which cooperative sequencing and

interactions give rise to an overall function at each moment of time (Luria, 1973). This widely accepted view of brain function as a complicated functional system became dominant in the 1950s and 1960s and is still the accepted view today. For example, since the 1980s new technologies such as functional MRI (fMRI), PET, SPECT and qEEG/MEG have provided ample evidence for distributed functional systems involved in perception, memory, drives, emotions, voluntary and involuntary movements, executive functions and various psychiatric and psychological dysfunctions (Mesulam, 2000). Modern PET, qEEG, MEG and fMRI studies are consistent with the historical view of “functional systems” presented by Luria in the 1950s (Luria 1973), i.e., there is no absolute functional localization because a functional system of dynamically coupled sub-regions of the brain is operating. For example, several fMRI and MRI studies (e.g., diffusion tensor imaging or DTI) have shown that the brain is organized by a relatively small subset of “Modules” and “Hubs” which represent clusters of neurons with high within cluster connectivity and sparse long distance connectivity (Hagmann et al, 2009; Chen et al, 2008; He et al, 2009). Modular organization is a common property of complex systems and ‘Small-World’ models in which maximum efficiency is achieved when local clusters of neurons rely on a small set of long distance connections in order to minimize the “expense” of wiring by shortened time delays between modules (Buzsaki, 2006; He et al, 2009). Also, recent qEEG and MEG analyses have demonstrated that important visually invisible processes such as directed coherence, phase delays, phase locking and phase shifting of different frequencies is critical in cognitive functions and various clinical disorders (Buzsaki, 2006; Sauseng and Klimesch, 2008; Thatcher et al, 2009a). Phase shift and phase synchrony has been shown to be one of the fundamental processes involved in the coordination of neural activity located in spatially distributed “modules” at each moment of time (Freeman and Rogers, 2002; Freeman et al, 2003; Sauseng and Klemish, 2008; Breakspear and Terry, 2002; Lachaux et al, 2000; Thatcher et al, 2005c; 2009; 2008b).

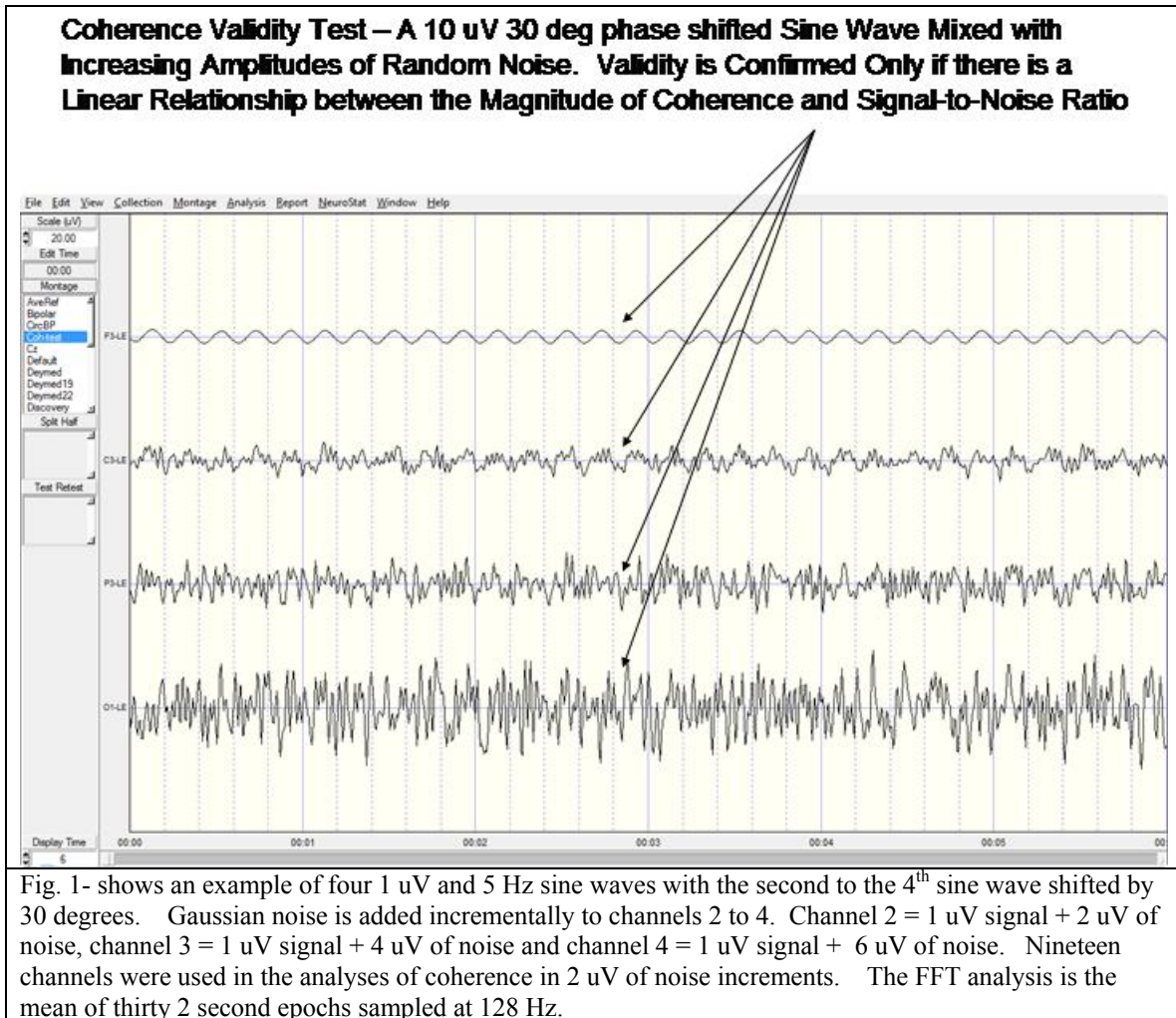
Validity of Coherence and Phase

Coherence is a measure of the stability of phase differences between two time series. Coherence is not a direct measure of an attribute like “temperature” or “volts”,

instead it is a measure of the “reliability” phase differences in a time series. If the phase differences are constant and unchanging over time then coherence = 1. If, on the other hand, phase differences are changing over time and are random over time then coherence = 0 (i.e., unreliable over time). Therefore, coherence is not a straightforward analytical measure like absolute power, rather coherence depends on multiple time samples in order to compute a correlation coefficient in the frequency or time domains. The validity and reliability of coherence fundamentally depends on the number of time samples as well as the number of connections (N) and the strength of connections (S) in a network or $\text{Coherence} = N \times S$. Coherence is sensitive to the number and strength of connections and therefore as the number or strength of connections decreases then coherence decreases because it is a valid network measure and as one would expect, the reliability of coherence declines when the number or strength of connections declines. Here is an instance where the validity of coherence is established by the fact that the reliability is low, i.e., no connections means no coupling and coherence approximates zero.

In order to evaluate the validity of coherence it is important to employ simulations using calibrated sine waves mixed with noise. In this manner a linear relationship between the magnitude of coherence and the magnitude of the signal-to-noise ratio can be demonstrated which is a direct measure of the predictive validity and concurrent validity of coherence and such a test is essential in order to evaluate the meaning of the reliability of coherence. For example, if one were to use an invalid method to compute coherence such as with an average reference, then it is irrelevant what the stability of the measure is because coherence is no longer measuring phase stability between two time series and therefore has limited physiological validity.

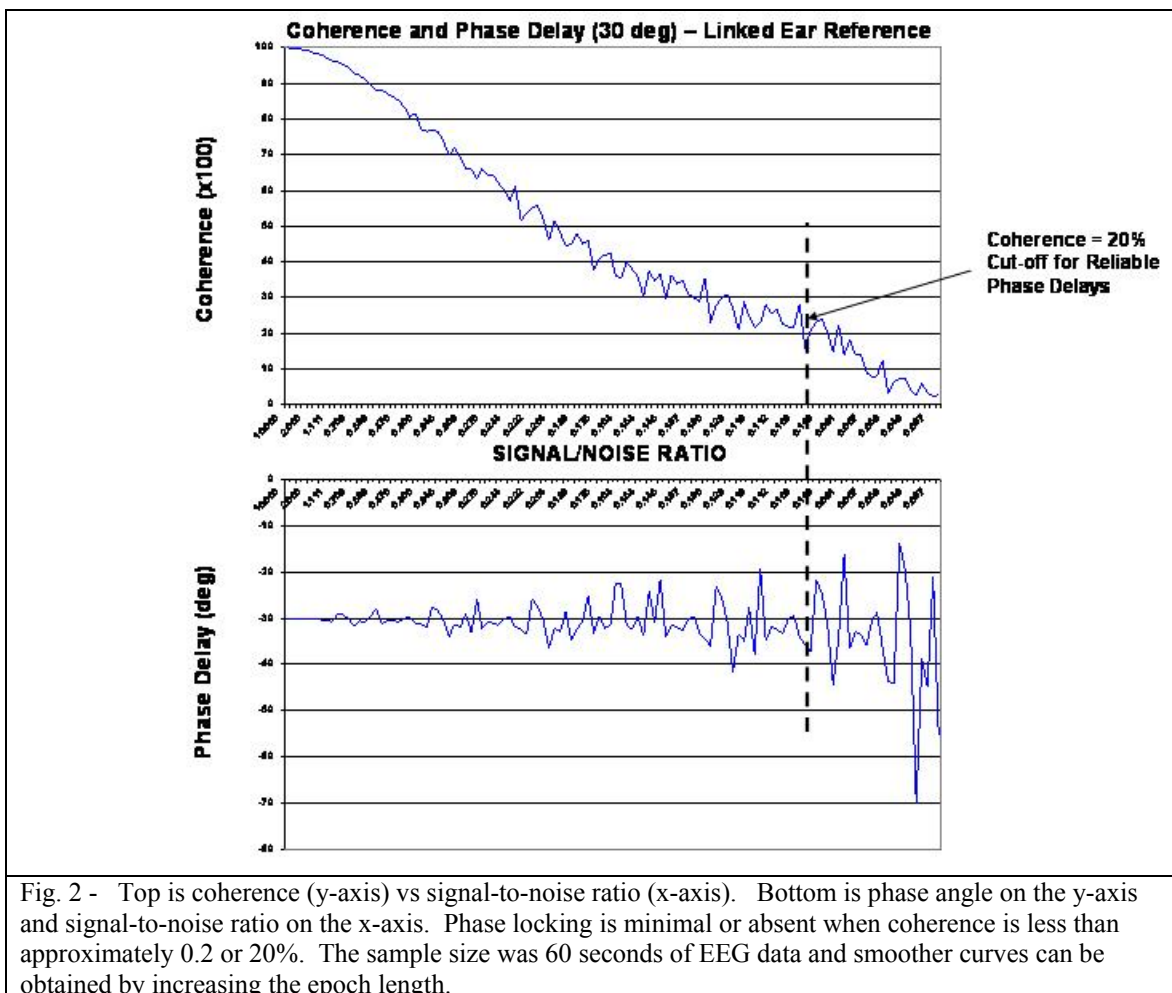
Figure 1 is an example of a validation test of coherence using 5 Hz sine waves and a 30 degree shift in phase angle with step by step addition of random noise. As shown in figure 1, a simple validity test of coherence is to use a signal generator to create a calibrated 1 uV sine wave at 5 Hz as a reference signal, and then compute



coherence to the same 1 uV sine wave at 5 Hz but shifted by 30 degrees and adding 2 uV of random noise, then the next channel add 4 uV of random noise, then 6 uV, etc.

Mathematically, validity equals a linear relationship between the magnitude of coherence and the signal-to-noise ratio, i.e., the greater the noise then the lower is coherence. If one fails to obtain a linear relationship then the method of computing coherence is invalid. If one reliably produces the same set of numbers but a non-linear relationship (i.e., no straight line) occurs then this means that the method of computing coherence is invalid (the method reliably produces the wrong results or is reliably off the mark).

Figure 2, shows the results of the coherence test in figure 1 that demonstrates a linear relationship between coherence and the signal-to-noise ratio, thus demonstrating that a standard FFT method of calculating coherence using a single common reference (e.g., one ear, linked ears, Cz, etc.) is valid. Note that the phase difference of 30 degrees



is preserved even when coherence is < 0.2 . The preservation of the phase difference and the linear decrease as a function of noise is a mathematical test of the validity of coherence.

Why the average Reference or Laplacian are Physiologically Invalid when Computing Coherence and Phase Differences

An important lesson in reliability and validity is taught when examining any study that fails to use a common reference when computing coherence. For example, the average reference mathematically adds the phase differences between all combinations of scalp EEG time series and then divides by the number of electrodes to form an average and then the average is subtracted time point by time point from the original time series recorded from each individual electrode thereby replacing the original time series with a

distorted time series. This process scrambles up the physiological phase differences so that they are irretrievably lost and can never be recovered. The method of mixing phase differences precludes meaningful physiological or clinical correlations since measures such as conduction velocity or synaptic rise or fall times can no longer be estimated due to the average reference. Also, coherence methods such as “Directed Coherence” can not be computed and more sophisticated analyses such as phase reset and phase shift and phase lock are precluded when using an average reference. The mixing together of phase differences in the EEG traces is also a problem when using the Laplacian transform and, similarly, reconstruction of EEG time series using Independent Component Analyses (ICA), also replaces the original time series with an altered time series that eliminates any physiological phase relationships and therefore is an invalid method of calculating coherence. One may obtain high reliability in test re-test measures of coherence, however, the reliability is irrelevant because the method of computation using an average reference or a Laplacian to compute coherence is invalid in the first place.

As pointed out by Nunez (1981) “The average reference method of EEG recording requires considerable caution in the interpretation of the resulting record” (p. 194) and that “The phase relationship between two electrodes is also ambiguous: (p. 195). As mentioned previously, when coherence is near unity then the oscillators are synchronized and phase and frequency locked. This means that when coherence is too low, e.g., < 0.2 , then the estimate of the average phase angle may not be stable and phase relationships could be non-linear and not synchronized or phase locked.

The distortions and invalidity of the average reference and Laplacian transform are easy to demonstrate using calibrated sine waves mixed with noise just as was done in figures 1 and 2. For example, figure three is the same simulation with a 30° phase shift as used for coherence with a common reference as shown in figure 2. The top row is coherence on the y-axis and the bottom row is the phase difference, the left column is using the average reference and the right column is the Laplacian. It can be seen in figure 3 that coherence is extremely variable and does not decrease as a linear function of signal-to-noise ratio using either the average reference nor the Laplacian montage. It can

also be seen in figure 3 that EEG phase differences never approximate 30 degrees and are extremely variable at all levels of the signal-to-noise ratio.

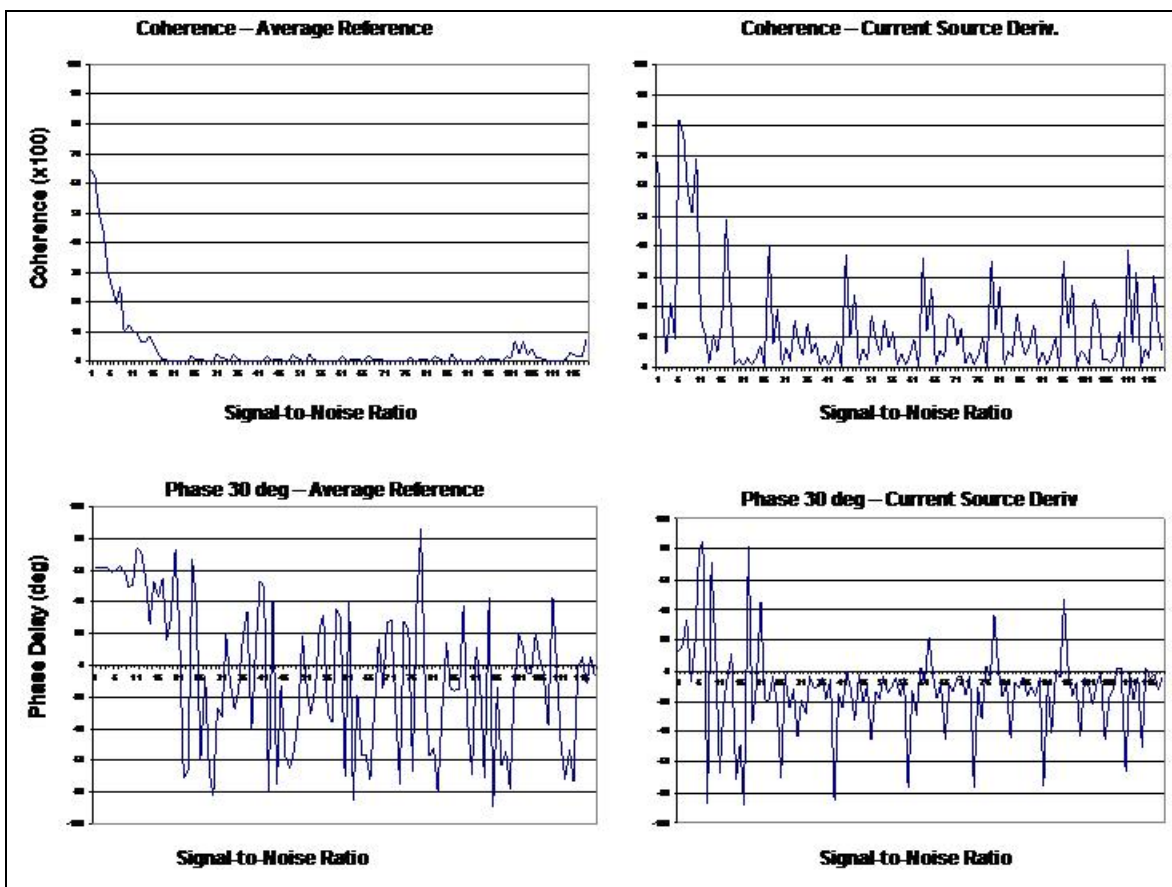


Fig. 3-. Left top is coherence (y-axis) vs signal-to-noise ratio (x-axis) with a 30^0 phase shift as shown in figure 2 using the average reference. The left bottom is phase differences in degrees in the y-axis and the x-axis is the signal-to-noise ratio using the average reference. The right top graph is coherence (y-axis) vs signal-to-noise ratio (x-axis) using the Laplacian montage. The right bottom is phase difference on the y-axis and signal-to-noise on the x-axis using the Laplacian montage. In both instances, coherence drops off rapidly and is invalid with no linear relationship between signal and noise. The bottom graphs show that both the average reference and the Laplacian montage fails to track the 30^0 phase shift that was present in the original time series. In fact, the phase difference is totally absent and unrepresented when using an average reference or a Laplacian montage and these simulations demonstrate that the average reference and the Laplacian montage are not physiologically valid because they do not preserve phase differences or the essential time differences on which the brain operates.

The results of these analyses are consistent with those by Rappelsberger, 1989 who emphasized the value and validity of using a single reference and linked ears in estimating the magnitude of shared or coupled activity between two scalp electrodes. The use of re-montage methods such as the average reference and Laplacian source derivation are useful in helping to determine the location of the sources of EEG of

different amplitudes at different locations. However, the results of this analysis which again confirm the findings of Rappelsberger, 1989 showed that coherence is invalid when using either an average reference or the Laplacian source derivation. This same conclusion was also demonstrated by Korzeniewska, et al (2003) and Essl and Rappelsburger (1998); Kamiński and Blinowska (1991); Kamiński et al (1997).

The average reference and the Laplacian transform also distort measures of phase differences which is also easy to demonstrate by using calibrated sine waves. For example, a sine wave at Fp1 of 5 Hz and 100 uV with zero phase shift, Fp2 of 5 Hz and 100 uV with 20 deg phase shift; F3 of 5 Hz and 100 uV with 40 deg phase shift; F4 of 5 Hz and 100 uV with 60 deg phase shift; C3 of 5 Hz and 100 uV with 80 deg phase shift; C4 of 5 Hz and 100 uV with 100 deg phase shift; P3 of 5 Hz and 100 uV with 120 deg phase shift; P4 of 5 Hz and 100 uV with 140 deg phase shift; O1 of 5 Hz and 100 uV with 160 deg phase shift and O2 of 5 Hz and 100 uV with 180 deg phase shift and channels F8 to Pz = 0 uV and zero phase shift. Figure 4 below compares the incremental phase shift with respect to Fp1 using Linked Ears common reference (solid black line), the Average Reference (long dashed line), and the Laplacian (short dashed line). This is another demonstration of how a non-common reference like the average reference and the Laplacian scramble phase differences and therefore caution should be used and only a common reference recording (any common reference and not just linked ears) is the only valid method of relating phase differences to the underlying neurophysiology, e.g., conduction velocities, synaptic rise times, directed coherence, phase reset, etc.

The analyses of the average reference and Laplacian to compute coherence should not be interpreted as a blanket statement that all of qEEG is invalid. On the contrary, when quantitative methods are properly applied and links to the underlying neuroanatomy and neurophysiology are maintained then qEEG analyses are highly reliable and physiologically valid. The lesson is that users of this technology must be trained and the use of calibration sine analyses should be readily available so that the users of qEEG can test basic assumptions themselves.

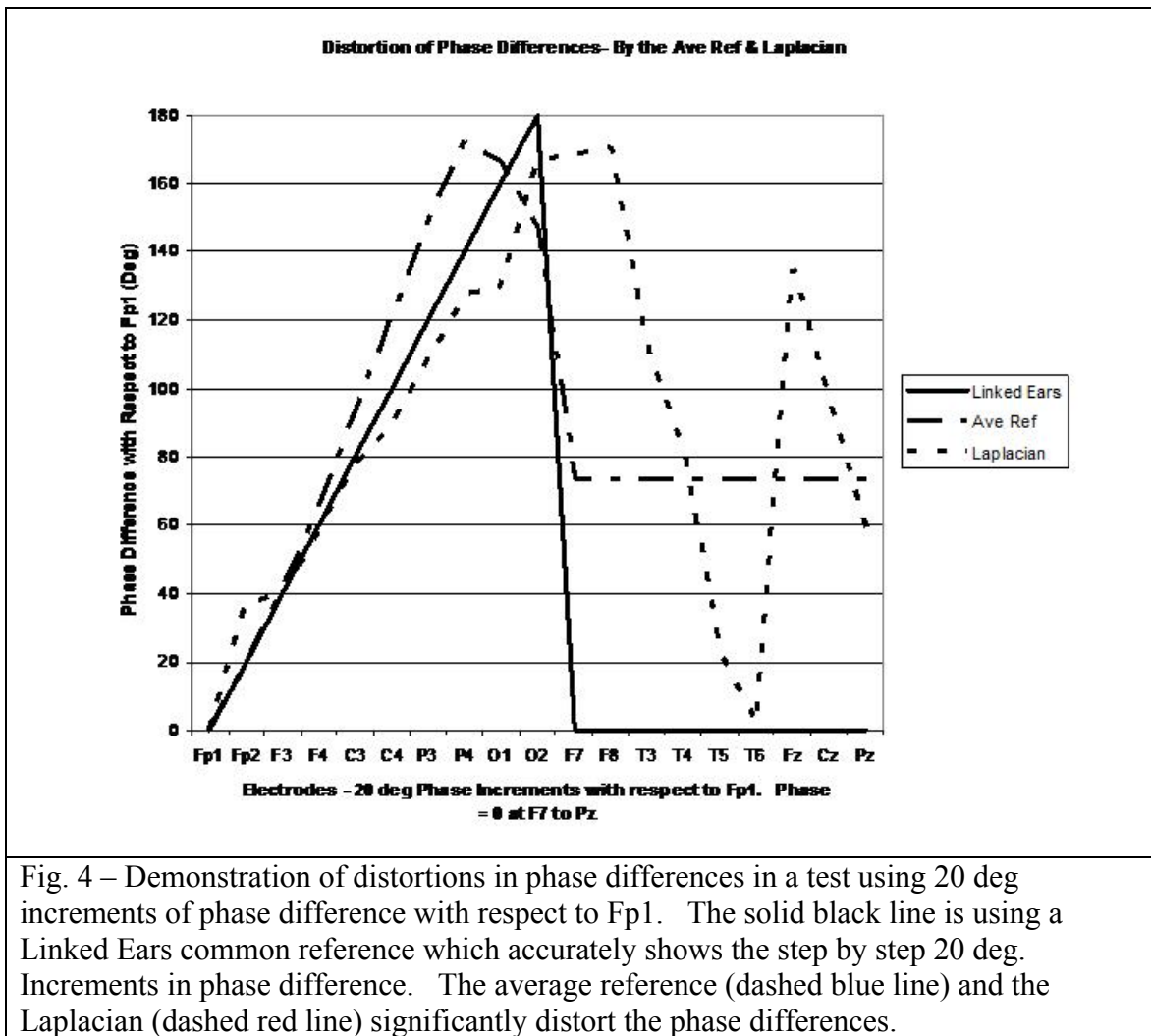


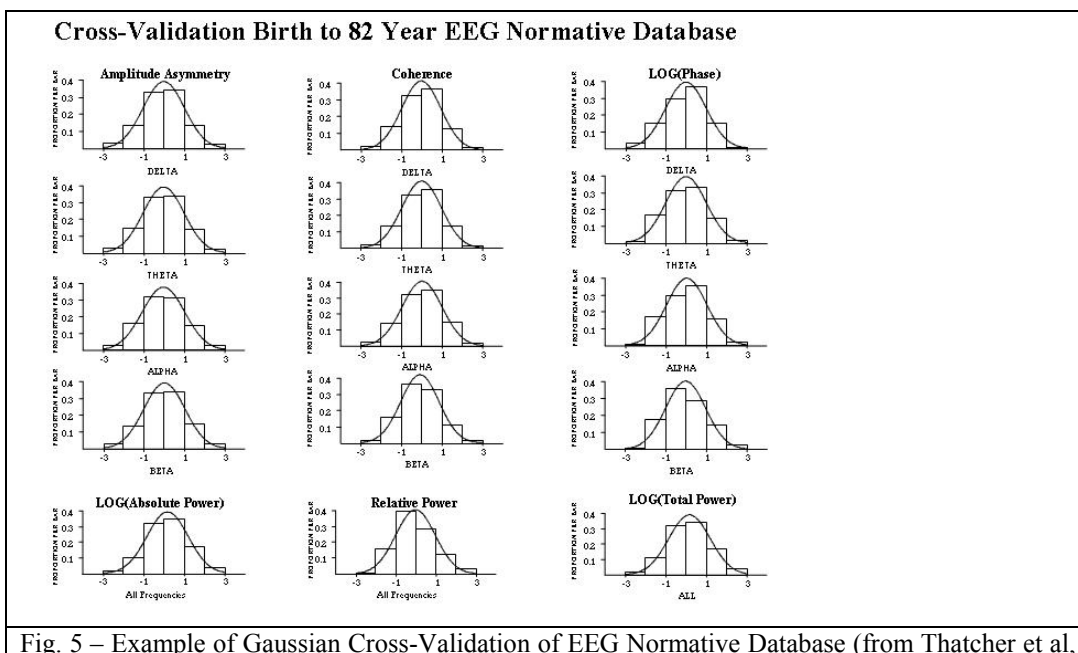
Fig. 4 – Demonstration of distortions in phase differences in a test using 20 deg increments of phase difference with respect to Fp1. The solid black line is using a Linked Ears common reference which accurately shows the step by step 20 deg. Increments in phase difference. The average reference (dashed blue line) and the Laplacian (dashed red line) significantly distort the phase differences.

Validity by Hypothesis Testing and qEEG Normative Data Bases

The Gaussian or Normal distribution is an ideal bell shaped curve that provides a probability distribution which is symmetrical about its mean. Skewness and kurtosis are measures of the symmetry and peakedness, respectively of the gaussian distribution. In the ideal case of the Gaussian distribution skewness and kurtosis = 0. In the real world of data sampling distributions skewness and kurtosis = 0 is never achieved and, therefore, some reasonable standard of deviation from the ideal is needed in order to determine the approximation of a distribution to Gaussian. The primary reason to approximate "Normality" of a distribution of EEG measures is that the sensitivity (i.e., true positive rate) of any normative EEG database is determined directly by the shape of the sampling

distribution. In a normal distribution, for example, one would expect that approximately 5% of the samples will be equal to or greater than ± 2 standard deviations and approximately 0.13 % ± 3 SD. (Hayes, 1973; John, 1977; John et al, 1987; Prichep, 2005; Thatcher et al, 2003a; 2003b).

A practical test of the sensitivity and accuracy of a database can be provided by cross-validation. There are many different ways to cross-validate a database. One is to obtain independent samples and another is to use a leave-one-out cross-validation method to compute Z scores for each individual subject in the database. The former is generally not possible because it requires sampling large numbers of additional subjects who have been carefully screened for clinical normality without a history of problems in school, etc. The second method is certainly possible for any database. Gaussian cross-validation of the EEG database used to evaluate TBI was accomplished by the latter method in which a subject is removed from the distribution and the Z scores computed for all variables based on his/her respective age matched mean and SD in the normative database. The subject is placed back in the distribution and then the next subject is removed and a Z score is computed and this process is repeated for each normal subject to obtain an estimate of the false positive hit rate. A distribution of Z scores for each of the EEG variables for each subject was then tabulated. Figure 5 is an example of the Gaussian



2003).

distributions of the cross-validated Z scores of 625 subjects from birth to 82 years of age used in a normative EEG database (Thatcher et al, 2003a).

Table I: Cross Validation of EEG Normative Database (from Thatcher et al, 2003).

Measure	% >2 SD	% <2 SD	% >3 SD	% <3 SD
Delta Amplitude Asym.	2.58	3.08	0.21	0.19
Theta Amplitude Asym.	2.29	2.62	0.15	0.13
Alpha Amplitude Asym.	2.71	2.72	0.18	0.19
Beta Amplitude Asym.	2.68	2.65	0.15	0.15
Delta Coherence	1.99	2.14	0.14	0.22
Theta Coherence	2.22	1.88	0.22	0.16
Alpha Coherence	2.55	1.62	0.18	0.18
Beta Coherence	2.20	1.38	0.18	0.10
Delta Phase †	0.89	3.52	0	0.23
Theta Phase †	1.61	1.87	0.04	0.13
Alpha Phase †	1.61	1.66	0.04	0.24
Beta Phase †	2.83	0.72	0.27	0.03
Absolute Power †	4.15	1.67	0.23	0.12
Relative Power	4.09	0.52	0.68	0
Total Power †	4.23	1.60	0.08	0.04
Average	2.58	1.98	0.18	0.14

† Data was logged transformed

Table I shows the results of a Gaussian cross-validation of the 625 subjects in the normative EEG database used in the evaluation of patients (Thatcher et al, 2003). A perfect cross-validation would be 2.3% at + 2 S.D., 2.3% at - 2 S.D., 0.13% at + 3 S.D. and 0.13 % at - 3 S.D. Table I shows a cross-validation grand average of $2.28\% \pm 2$ S.D. and $0.16\% \pm 3$ S.D. The cross-validation result shows that the EEG normative database is statistically accurate and sensitive with slight differences between variables that should be taken into account when evaluating individual Z scores.

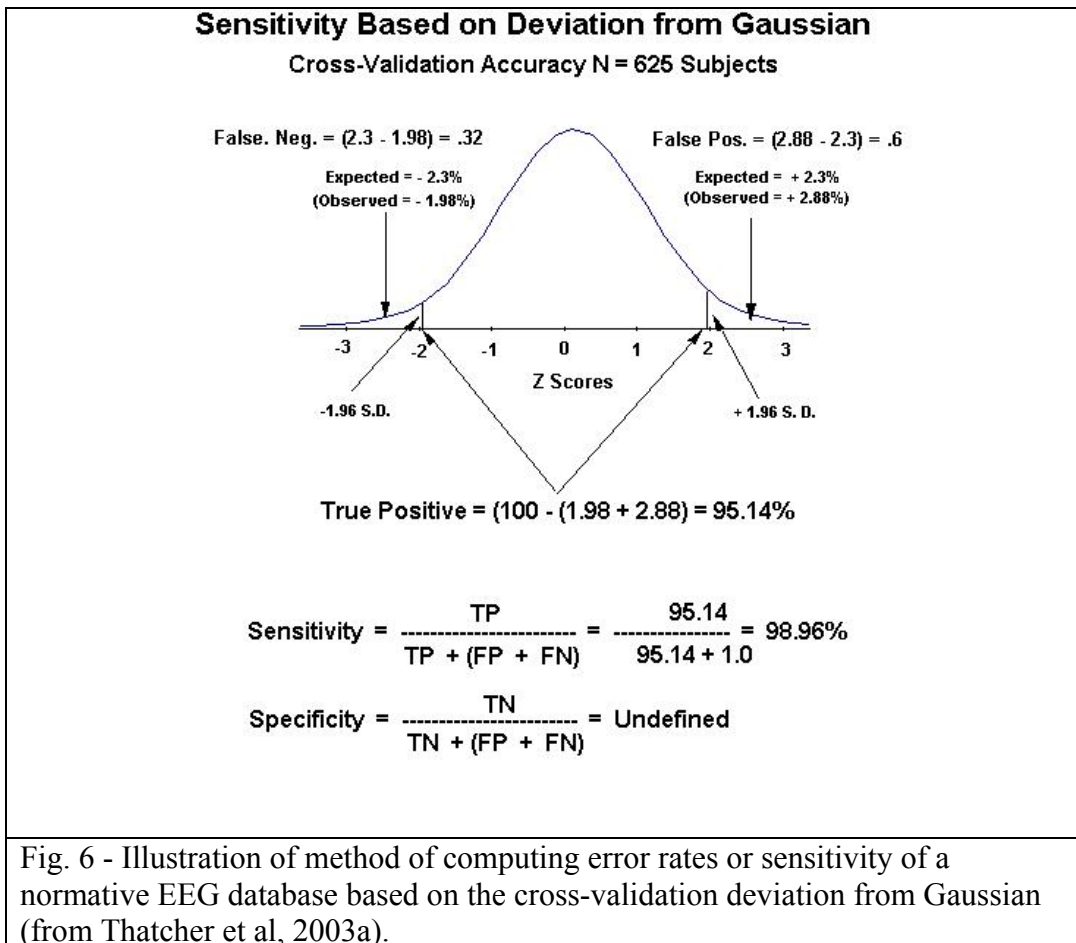


Figure 6 is a bell shaped curve showing the ideal Gaussian and the average cross-validation values of the EEG normative database used to evaluate patients. The error rates or the statistical sensitivity of a qEEG normative database are directly related to the deviation from a Gaussian distribution. Figure 6 also illustrates the method of estimating the statistical sensitivity of a normative EEG database in terms of the deviation from Gaussian.

Table II is an example of the calculated sensitivity of a EEG normative database for different age groups using the method described in figure 6.

FFT Normative Database Sensitivities				
2 STDEVs	CALC SENSITIVITY: $FP=TP/(TP+FP)$ or $FN=TP/(TP+FN)$			
AGES	(+/- 2 SD)	(>= 2 SD)	(<= -2 SD)	
0-5.99	0.95448265	0.9771774	0.97730526	+/- 2 Std. Dev.
6-9.99	0.95440363	0.9772031	0.97720054	
10-12.99	0.9543997	0.97724346	0.97715624	
13-15.99	0.95440512	0.97723601	0.97716911	
16-ADULT	0.9543945	0.97718143	0.97721307	
ALL	0.95442375	0.97720714	0.97721661	
3 STDEVs	CALC SENSITIVITY: $FP=TP/(TP+FP)$ or $FN=TP/(TP+FN)$			
AGES	(+/- 3 SD)	(>= 3 SD)	(<= -3 SD)	
0-5.99	0.99743898	0.99871123	0.99872774	+/- 3 Std. Dev.
6-9.99	0.99744112	0.99871611	0.99872501	
10-12.99	0.99744688	0.99873171	0.99871518	
13-15.99	0.99743186	0.99871951	0.99871234	
16-ADULT	0.99743835	0.99870216	0.99873619	
ALL	0.99744002	0.99871716	0.99872286	
Table II – Normative EEG database sensitivities for different age groups at +/- 2 standard deviations and +/- 3 standard deviations (from Thatcher et al, 2003a).				

Predictive Validity of Normative Databases

Predictive (or criterion) validity has a close relationship to hypothesis testing by subjecting the measure to a discriminant analysis or cluster analysis to some statistical analysis in order to separate a clinical sub-type from a normal reference database. Nunnally (1978) gives a useful definition of predictive validity as: “when the purpose is to use an instrument to estimate some important form of behavior that is external to the measuring instrument itself, the latter being referred to as criterion [predictive] validity.” For example, science “validates” the clinical usefulness of a measure by its false positive and false negative rates and by the extent to which there are statistically significant correlations to other clinical measures and, especially, to clinical outcomes (Hughes and John, 1999).

An example of predictive validity of the Linked Ears qEEG normative database is the use of a discriminant function to evaluate the false positive/false negative classification rate using a normative database and TBI patients (Thatcher et al, 1989). In

this study the traumatic brain injured patients were distinguished from age matched normal control subjects at a classification accuracy = 96.2% . Four different cross-validations were conducted in the Thatcher et al (1989) study and showed similar accuracies although the strength of the discrimination declined as a function of time from injury to test.

Correlation : QEEG and Neuropsychological Test Scores in TBI Patients. J. Neuropsychiatry & Clin. Neurosci., 13: 77-87, 2001

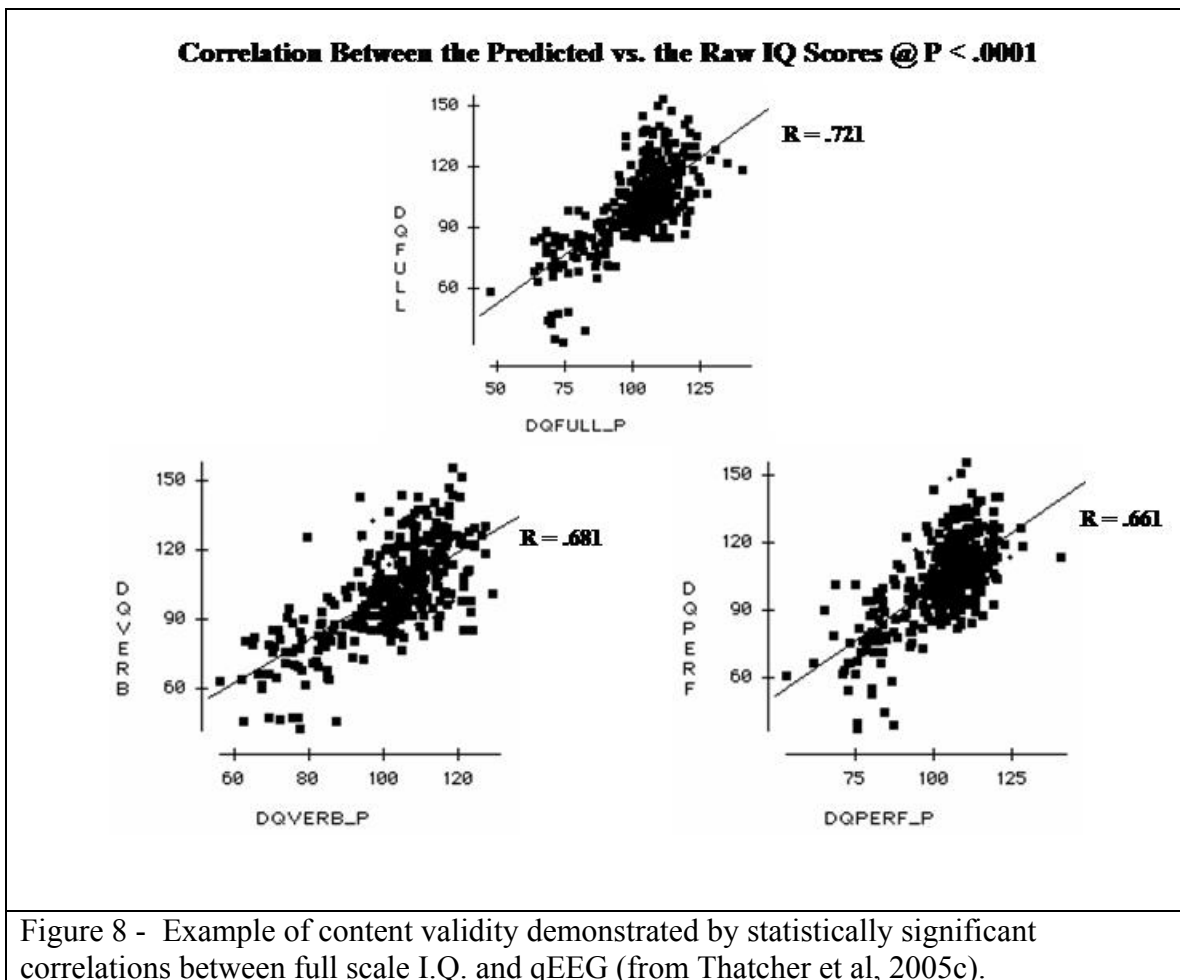
	Correlation	Probability
WAIS TEST-Scaled Scores		
Vocabulary	-0.416	0.05
Similarities	-0.640	0.001
Picture Arrangement	-0.576	0.01
Performance	-0.504	0.01
Digit Symbol	-0.524	0.01
ATTENTION TEST-Raw Scores		
Trail Making A-Response Time	0.627	0.001
Trail Making B-Response Time	0.627	0.001
Stroop-Word	-0.427	0.05
Stroop-Color	-0.618	0.001

Examples of Predictive Validity of QEEG in the Evaluation Of Traumatic Brain Injury

Fig. 7 – Example of predictive and content validity by clinical correlations of qEEG with Neuropsychological test scores (Thatcher et al, 2001a).

Figure 7 shows the correlation to neuropsychological test scores in an independent replication of the Thatcher et al (1989) study. In this study a similar discriminant function produced similar sensitivities and also predicted the Glasgow Coma Score with a correlation of 0.85 (Thatcher et al, 2001a). Another example of predictive validity is the ability of qEEG normative values to predict cognitive functioning. Figure 8 shows correlations to Full Scale I.Q. as an example of predictive validity and content validity . A more complete analysis of the predictive validity of a normative EEG database is shown in Table III (Thatcher et al, 2003; 2005a; 2005b). In Table III the percentage of statistically significant correlations at $P < .01$. between qEEG normative EEG and

WRAT School Achievement scores and measures of intelligence are shown. The relative effect size of the normative EEG correlations differs for different measures which is valuable information when using any normative database, not just a qEEG normative database. Similar high and significant correlations between qEEG and neuropsychological test performance have been published in many studies. A search of the National Library of Medicine's database using the search terms: EEG and Neuropsychological Tests produced 1,351 citations.



EFFECT SIZE P < .01: qEEG Correlations with School Achievement & IQ Measures						
Percent Significant Correlations @ P < .01, N = 466						
Amplitude Asymmetry						
P <= .01	READING	SPELLING	ARITH	IQFULL	IQVERB	IQPERF
DELTA	64%	61%	55%	64%	61%	61%
THETA	78%	70%	70%	70%	67%	59%
ALPHA	63%	63%	53%	64%	63%	52%
BETA	56%	56%	34%	58%	61%	47%
Coherence						
P <= .01	READING	SPELLING	ARITH	IQFULL	IQVERB	IQPERF
DELTA	27%	14%	41%	38%	22%	38%
THETA	27%	6%	36%	30%	27%	23%
ALPHA	9%	6%	45%	11%	14%	5%
BETA	11%	5%	38%	22%	17%	6%
Absolute Phase						
P <= .01	READING	SPELLING	ARITH	IQFULL	IQVERB	IQPERF
DELTA	11%	8%	8%	16%	6%	17%
THETA	9%	5%	8%	13%	9%	17%
ALPHA	9%	3%	33%	14%	19%	6%
BETA	9%	5%	30%	6%	9%	3%
Relative Power						
P <= .01	READING	SPELLING	ARITH	IQFULL	IQVERB	IQPERF
DELTA	13%	0%	31%	0%	6%	0%
THETA	56%	44%	94%	6%	6%	0%
ALPHA	19%	0%	75%	0%	0%	0%
BETA	13%	6%	44%	19%	13%	13%
Relative Power Ratios						
P <= .01	READING	SPELLING	ARITH	IQFULL	IQVERB	IQPERF
Theta/Beta	50%	44%	63%	56%	56%	50%
Theta/Alpha	13%	0%	69%	0%	0%	0%
Alpha/Beta	50%	31%	50%	38%	38%	25%
Delta/Theta	19%	25%	56%	19%	13%	25%

Table III – Examples of predictive validity by clinical correlations between qEEG and intelligence (WISC-R) and academic achievement tests (WRAT) (from Thatcher et al, 2003a).

Examples of Content Validity of Normative Databases

Content validity is defined by the extent to which an empirical measurement reflects a specific domain of content. For example, a test in arithmetic operations would not be content valid if the test problems focused only on addition, thus neglecting subtraction, multiplication and division. By the same token, a content-valid measure of

cognitive decline following a stroke should include measures of memory capacity, attention and executive function, etc.

Normative databases are distinct from small experimental control groups in their scope and their sampling restriction to clinically normal or otherwise healthy individuals for the purpose of comparison. Another distinguishing characteristic of normative databases is the ability to compare a single individual to a population of “normal” individuals in order to identify the measures that are deviant from normal and the magnitude of deviation. Normative databases themselves do not diagnose a patient’s clinical problem. Rather, a trained professional first evaluates the patient’s clinical history and clinical symptoms and complaints and then uses the results of normative database comparisons in order to aid in the development of an accurate clinical diagnosis. Most importantly to link functional localization of deregulated brain regions (i.e., anatomical hypotheses) to a patient’s symptoms and complaints.

There are many examples of the clinical content validity of qEEG and normal control groups in ADD, ADHD, Schizophrenia, Compulsive disorders, Depression, Epilepsy, TBI and a wide number of clinical groupings of patients as reviewed by Hughes and John, (1999). In most of these studies an assortment of clinical measures were correlated to a variety of brain EEG sources related to the disorder under study. One of the most consistent and relevant findings is anatomical localization related to different psychiatric and psychological disorders, e.g., cingulate gyrus and depression, right parietal lobe and spatial neglect, left angular gyrus and dyslexia, etc. qEEG anatomical correlations with clinical disorders form the foundation of modern day qEEG which is another example of content validity. Since 1999, the number of qEEG studies demonstrating anatomical and frequency clinical content validity is several hundred. For example, all clinical LORETA qEEG studies demonstrate anatomical content validity in that there are no published studies showing low localization accuracy when using LORETA. The term “Low Resolution Electromagnetic Tomography” refers to a “smearing” around the spatially accurate maximum in the center of a spatial volume. This is defined by the point-spread function of the Laplacian spatial operator in LORETA (Pascual-Marqui et al, 1994; Pascual-Marqui, 1999). This means that LORETA is spatially accurate but with a smeared resolution like a probability cloud. Clinical

correlations consistent with PET and SPECT and fMRI are abundant in today's scientific literature (see the National Library of Medicine database at: <https://www.ncbi.nlm.nih.gov/sites/entrez> and see the section on "Validity of LORETA" for some specific citations.

Anatomical Hypothesis Testing and Planned qEEG Comparisons

The best use of parametric statistics is to form hypotheses prior to conducting an analysis in a procedure referred to as "Planned Comparisons" (Hayes, 1973). In this manner, one does not need to resort to multiple comparisons which are performed only when an experimenter has no idea about what the test is likely to yield and is totally ignorant of possible statistically significant differences. Because one has no idea what to expect it is not possible to form hypotheses and one then must resort to multiple comparisons which have high Type II errors (saying something is false when it is not false) in order to reduce the Type I errors (saying something is true when it is not true) because of the total ignorance of possible relationships between groups or between variables.

Planned comparisons are more robust and valid than multiple comparisons because specific hypotheses are generated prior to conducting statistical tests which markedly minimizes the probability of both Type I and Type II errors. A complaint against qEEG is that there are such a large number of statistical tests and one would expect 5% to be significant by chance alone. The problem with this argument is that the 5% by chance must be random in space and in qEEG features. The random chance argument falls to the way side when there are focal anatomical deviations that were predicted prior to analysis. Additional content validity is when the deviant qEEG findings are located in anatomical regions known to be linked to the patient's symptoms and clinical history. For example, the MRI uses approximately 10,000 voxels and one would expect 500 to be significant by chance at $P < .05$ if these 500 voxels are randomly distributed throughout the volume. However, if 100 voxels are statistically significant in the right parietal lobe which happens to be where the patient was struck on the head, then the 5% significant multiple test argument is not valid and must be discarded. The same is true for the qEEG, for example, if one uses planned comparisons and predicts that the

left parietal lobe will be deviant from normal in a dyslexic child prior to recording EEG and the qEEG shows many deviations from normal in the left parietal region then this can not be explained by chance alone. The use of planned comparisons is especially useful when using LORETA source localization methods because thousands of voxels are involved. An example, of planned comparisons is in figure 9. Here the surface qEEG analyses showed focal deviation from normal in the right hemisphere in a patient that was struck with a bat near to his right parietal lobe. The sources of the right parietal lobe deviations from normal are then predicted to appear in particular Brodmann areas prior to launching LORETA. Once LORETA is launched then the frequency and anatomical hypotheses can be tested to determine their accuracy and validity.

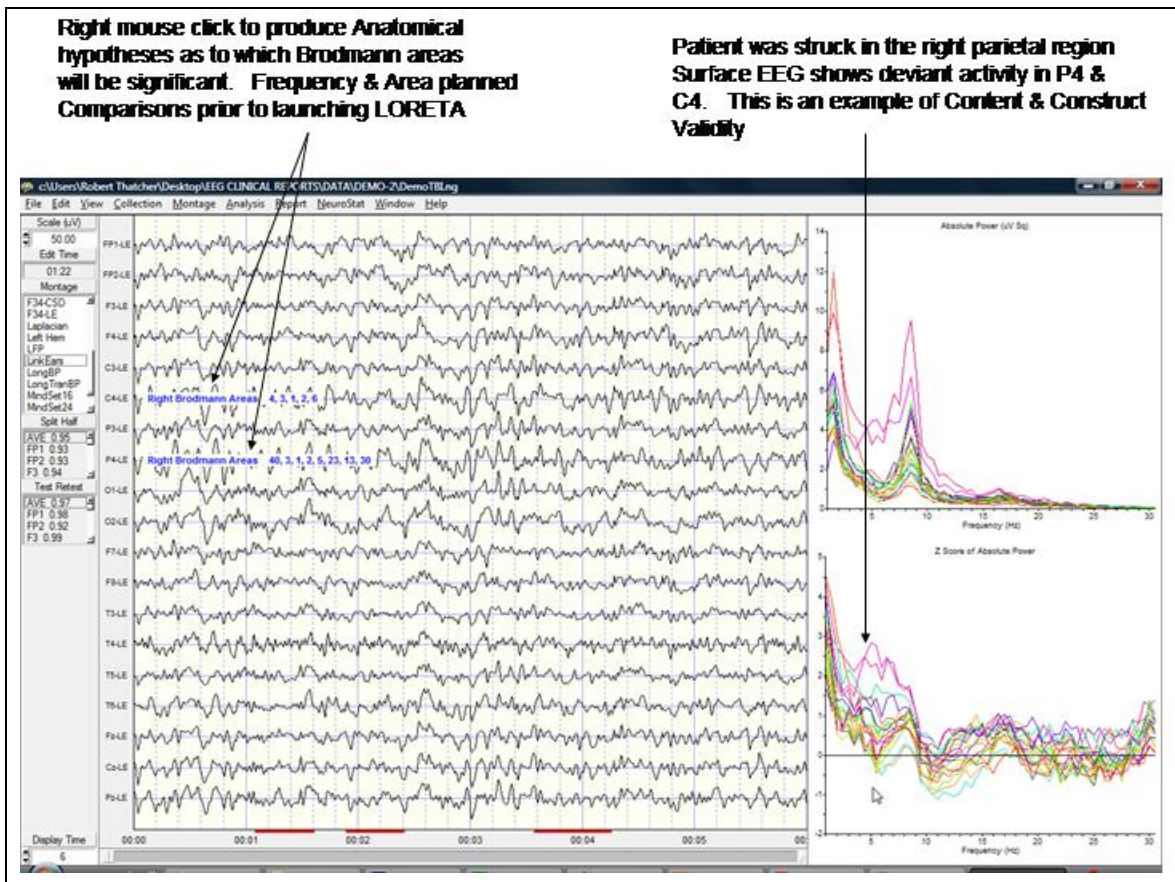


Fig. 9- Example of “Planned comparisons” using hypothesis creation prior to launching LORETA. Content and construct validity are present because the patient was hit on the right parietal lobe and the right parietal lobe shows deviant EEG activity (e.g., > 2 st. dev.) Further construct validity is established by LORETA analyses that confirm anatomical hypotheses based on the surface EEG locations and frequencies of deviance.

Predictive Validity and qEEG

Predictive validity is sometimes referred to as “criterion validity” and has a close relationship to hypothesis testing by subjecting the measure to an independent test of its ability to predict clinical measures such as severity of injury or intelligence, attention, executive function, etc. Nunnally (1978) gives a useful definition of predictive validity as: “when the purpose is to use an instrument to estimate some important form of behavior that is external to the measuring instrument itself, the latter being referred to as criterion-validity.” For example, one “validates” a written driver’s license test by hypothesizing that it accurately predicts how well some group of persons can operate an automobile. If the driving test fails to predict driving competence, then the test must be rejected or replaced. In the case of traumatic brain injury (TBI) one “validates” the qEEG by showing that it accurately predicts severity of TBI as measured by Hospital admission scores such as the Glasgow Coma Score (GCS) or length of coma or in other independent tests such as neuropsychological tests, etc. (Hughes and John, 1999).

False Positive and False Negative Error Rates of qEEG: Example of Content Validity in Traumatic Brain Injury

Peer reviewed scientific publications of 608 mild TBI patients compared to 108 age matched normal subjects demonstrated, in independent cross-validations an average false positive rate approximately 5% and an average false negative rate of approximately = 10% to 15% (Thatcher et al, 1989). Similar levels of sensitivity (the probability that a test result will be positive when the disorder is present) and specificity (the probability that at test result will be negative when the disorder is not present) were reported in a series of independent and replicated qEEG studies of TBI for the detection of a pattern consistent with traumatic brain injury as a causal agent (Thatcher et al, 1991; 2001a; Thornton, 1999; Thornton and Cormody, 2005 and Leon-Carrion et al, 2008a). Obtaining a content-valid measure of any phenomena involves at least three interrelated steps: 1- one must be able to specify the full domain of content that is relevant, 2- one must be able to identify the selection of relevant measures from the larger universe of possible measures with the understanding that over sampling is usually necessary and 3-

one must be able to test the content validity of the measuring instrument and/or be able to cite the peer reviewed literature in which the content-validity of the qEEG had been tested. As stated by Cronback (1977, pg. 447) “One validates, not a test, but an *interpretation of data arising from a specified procedure*”. This distinction is crucial because it is quite possible for a measuring instrument to be relatively valid for measuring one kind of phenomenon but entirely invalid for assessing other phenomena. The purpose of qEEG discriminant functions is not to derive a diagnosis because the diagnosis should be based on the patient’s clinical history and symptoms and complaints. qEEG discriminant functions are designed to further evaluate the extent, locations and severity of the EEG patterns that are present in individuals already diagnosed with a disorder.

qEEG involves the measurement of a relatively large number of electrical processes some of which may be affected by a traumatic brain injury (TBI). For example, animal studies and imaging studies in humans have demonstrated that maximal damage to the brain following TBI occurs at the interface between the brain and the skull bone (Ommaya, 1968; 1971; 1995). Another primary and common injury to the brain due to TBI are “shear” forces in which rapid acceleration/deceleration results in different brain parts moving at different rates, for example, the gray matter moves faster and further than the white matter thus stretching axonal fibers, etc. (Ommaya, 1968). Thus, a content valid qEEG measure of TBI should be capable of measuring electrical activity in frontal and temporal lobes where the brain-to-skull forces are greatest. Similarly, a content valid qEEG test of TBI must be capable of measuring EEG phase and EEG coherence which reflect the axonal conduction velocities and long distance cortical communication linkages (Thatcher et al, 1989; 1998b; 2001). If these measures are omitted then the test is not valid for the same reason that a test of arithmetic is invalid if it omits addition and subtraction. Over the years there is reasonable consistency of qEEG findings in TBI across studies which can be summarized by: 1- reduced power in the higher frequency bands (8 to 40 Hz) which is linearly related to the magnitude of injury to cortical gray matter, 2- increased slow waves in the delta frequency band (1 to 4 Hz) in the more severe cases of TBI which is linearly related to the magnitude of cerebral white matter injury and, 3- changes in EEG coherence and EEG phase delays which are linearly

related to the magnitude of injury to both the gray matter and the white matter, especially in frontal and temporal lobes (Thatcher, 2008).

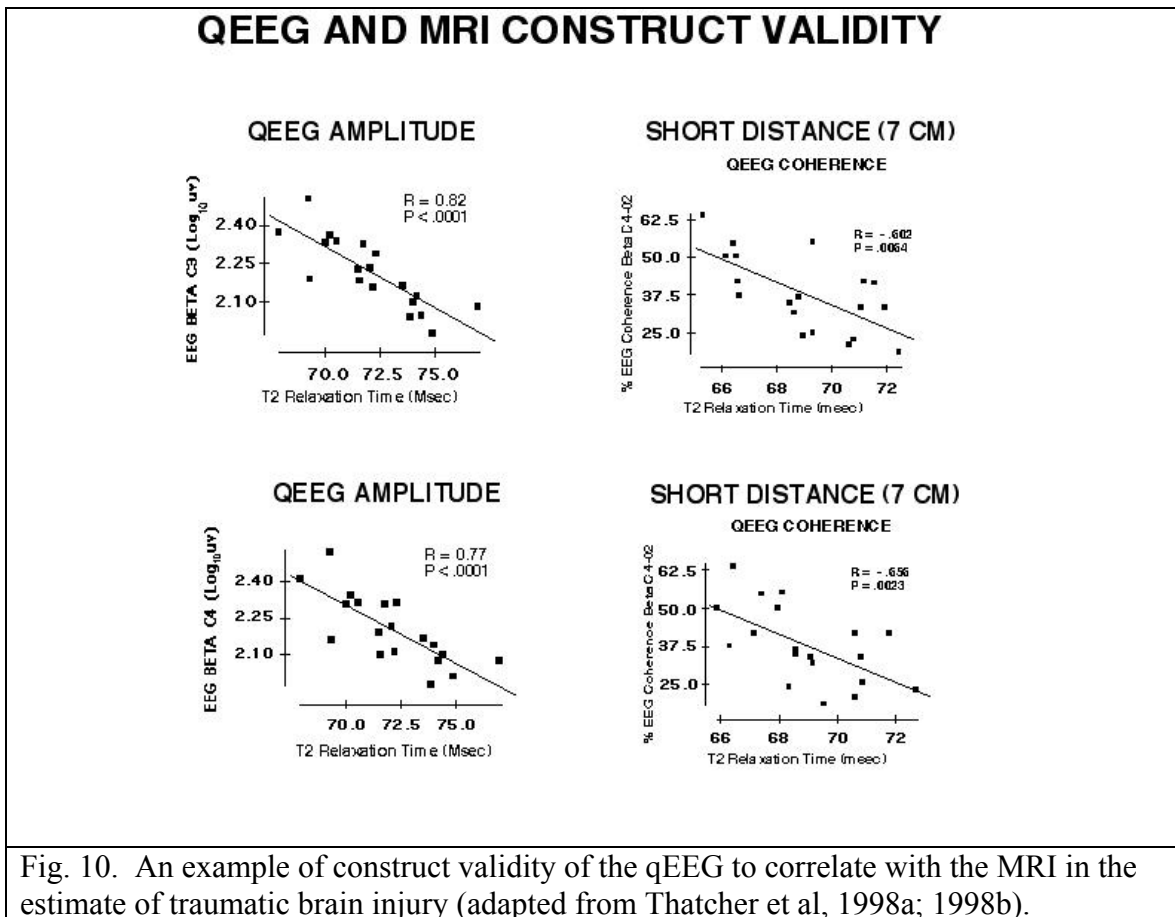
qEEG Construct Validity

Construct validity is concerned with the validity of empirical measures and hypothesis testing of theoretical concepts. As Carmines and Zeller (1979) state: “Construct validity is concerned with the extent to which a particular measure relates to other measures consistent with theoretically derived hypotheses concerning the concepts that are being measured”. Construct validity typically involves three steps: 1- the theoretical relationship between the concepts themselves must be specified and testable hypotheses stated, 2- the empirical relationship between the measures of the concepts must be examined and, 3- the empirical evidence must be interpreted in terms of how it affirms, rejects or clarifies the construct validity of the particular measure.

For example, in qEEG measures of traumatic brain injury one hypothesis is that rapid acceleration/deceleration contuses (bruises) brain tissue especially where the brain sits on the bony skull vault (Ommaya, 1968; 1995), another theory is that damage to neuronal membranes will result in reduced ionic flows and reduced amplitude of the EEG and high frequencies and a shift in frequency toward the theta and delta frequencies (lower frequency ranges). These two theoretical hypotheses regarding which qEEG measures would be expected to change following TBI have been tested and confirmed in the peer reviewed scientific literature (Randolph and Miller, 1998; Thatcher et al, 1989; 1991; 2001; 1998a; 1998b; Thornton, 1999; Thornton and Carmody, 2005; Leon-Carrion, 2008a; 2008b; Cao et al, 2008).

The qEEG is also used for prognoses in the neurointensive care unit. Fabregas et al (2004) reported a cross-validation performance error of 3.06% (95% confidence interval) for predicting recovery from coma. Similar accuracy of predicting recovery of consciousness was reported by others (Shields et al, 2007; Buzea, 1995; Jordan, 1993; Scheuer, 2002, Claassen, 2000; Hyllienmark and Amark, 2007; Kane et al, 1998; Thatcher et al, 1991). Jordan (1993) reported that qEEG can impact medical decision-making in 81% of the monitored patients and Claassen et al (2000) reported that qEEG findings influenced therapeutic management with decisive decisions on many occasions.

Figure 10 is an example of construct validity of the qEEG in the measurement of TBI in which correlations of MRI were used to test the null hypothesis = 0, about damage to the average concentration of ionic channels in a volume of cortex that produces EEG (Thatcher et al, 1998a; 1998b; 2001b).



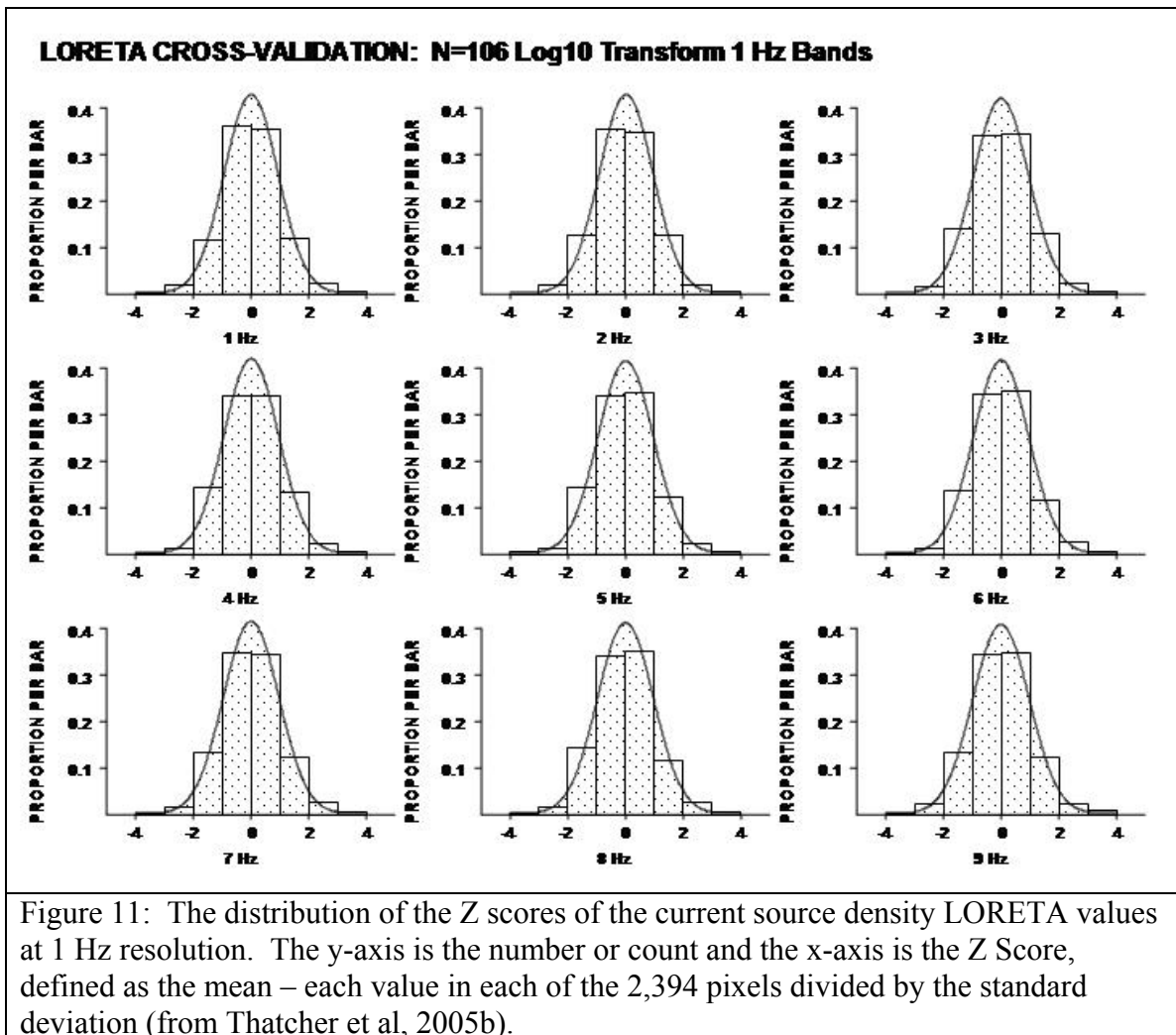
In fig. 10, construct validity of qEEG was tested by examining the hypothesized relationship between the integrity of gray matter membranes using the MRI and the amplitude and coherence of the EEG. The hypothesis predicted reduced connectivity and a decline in amplitude of the EEG related to reduced integrity of neural membranes. The results of the construct validity tests of the qEEG in TBI were born out as valid as reported in peer reviewed publications (Thatcher et al, 1998a; 1998b; 2001b). These same studies also tested content validity by correlating the independent MRI measures with selected qEEG measures and finally, predictive validity was also tested by correlations with neuropsychological test scores which co-varied with both the qEEG and

the MRI in a predictable manner. A similar cross-validation study was performed by Korn et al, (2005) showing significant correlations between LORETA current source activity and SPECT scans in TBI patients.

Validity of a LORETA qEEG Normative Database

There are over 795 peer reviewed journal articles on the use of LORETA for the identification of the 3-dimensional sources of the EEG in many different clinical populations. Because different regions in the brain are involved in different functional systems, then the reliability and validity of LORETA is established by the degree to which accurate localization is demonstrated and by repeatability across subjects and across experiments. It is easy to demonstrate that different samples of EEG yield the same localization and/or that a particular local event in the EEG corresponds to an expected source of that event, for example, alpha spindles maximum in O1 and O2 are localized to the occipital cortex by LORETA and not some where unexpected, e.g., right temporal lobe, etc. This is an example of content validity.

The reliability and validity of LORETA source localization can be demonstrated using mathematical simulations and standard tests in Systat and SPSS as well as by determining that the distribution of current sources is represented by a Gaussian distribution. To the extent the individual variables are Gaussian distributed then the mathematics of parametric statistics are valid and useful. Thus, step one in evaluating the validity of a LORETA normative database is to test and establish that the current sources are Gaussian distributed. Figure 11 shows the distribution of current source densities after \log_{10} transform in 1 Hz frequency bands from 1 to 9 Hz. Figure 11 also shows that a reasonable approximation to a Gaussian distribution was achieved by the \log_{10} transform. The distribution of current source densities after the Box-Cox transform were essential the same as for the \log_{10} and therefore are not displayed.



Standard cross-validation methods can also be used to establish reliability and validity. That is, the classification of normal subjects as not being normal by a leave-one-out cross-validation procedure or by a direct cross-validation procedure provides an estimate of the false positives (Type I error) and false negatives (Type II error) of the normative database. Table IV shows the skewness and kurtosis of the \log_{10} transformed data and the percentages of Z scores at ± 2 standard deviations and ± 3 standard deviations for each of the 1 Hz frequency bands for the eyes closed condition for linked ears reference. The sensitivities ranged from 95.64% at 2 standard deviations to 99.75% at 3 standard deviations. Average skewness = 0.29 and average kurtosis = 0.68. Thus, gaussianity can be approximated at a frequency resolution of 1 Hz.

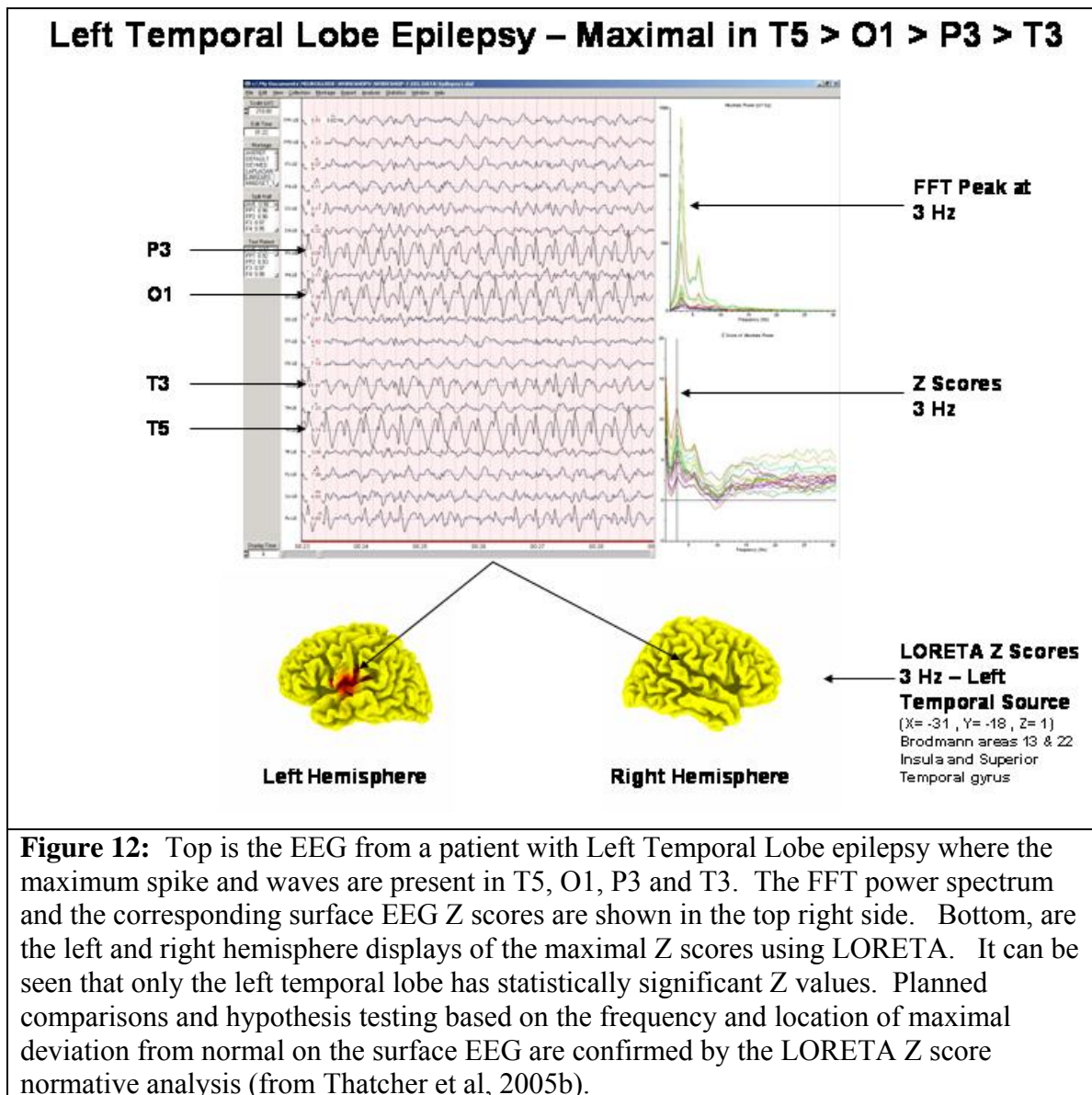
LORETA CROSS-VALIDATION: NORMs_LE_EC_ADULTs_z(LOG10(LORETA))_n=106										
			1 STANDARD DEVIATIONS		2 STANDARD DEVIATIONS		3 STANDARD DEVIATIONS			
Frequency	SKEWNESS	KURTOSIS	z < -1 SD	z > 1 SD	z < -2 SD	z > 2 SD	z < -3 SD	z > 3 SD	SEN 2SD	SEN 3SD
FREQ 1	0.08	0.84	13.76%	14.60%	2.22%	2.52%	0.51%	0.45%	95.64%	99.75%
FREQ 2	0.19	0.39	14.47%	15.20%	1.79%	2.44%	0.21%	0.46%	95.64%	99.75%
FREQ 3	0.24	0.14	15.79%	15.62%	1.66%	2.54%	0.03%	0.35%	95.64%	99.74%
FREQ 4	0.29	0.16	15.73%	15.83%	1.21%	2.63%	0.02%	0.42%	95.64%	99.74%
FREQ 5	0.27	0.17	15.79%	15.13%	1.44%	2.85%	0.00%	0.41%	95.64%	99.74%
FREQ 6	0.33	0.32	15.39%	14.77%	1.62%	3.11%	0.01%	0.54%	95.64%	99.75%
FREQ 7	0.33	0.30	15.25%	15.44%	1.79%	2.99%	0.00%	0.50%	95.64%	99.75%
FREQ 8	0.38	0.47	15.90%	14.69%	1.72%	3.19%	0.00%	0.44%	95.65%	99.75%
FREQ 9	0.21	0.29	15.66%	15.22%	2.20%	2.88%	0.00%	0.32%	95.65%	99.74%
FREQ 10	0.17	0.23	14.52%	16.43%	2.18%	2.38%	0.00%	0.30%	95.64%	99.74%
FREQ 11	0.64	1.51	12.62%	15.23%	1.65%	3.21%	0.00%	0.87%	95.64%	99.75%
FREQ 12	0.29	0.80	13.91%	15.06%	1.88%	3.17%	0.23%	0.44%	95.65%	99.75%
FREQ 13	0.22	0.82	13.97%	15.20%	1.83%	2.93%	0.44%	0.45%	95.64%	99.75%
FREQ 14	0.25	0.82	14.08%	14.75%	1.79%	3.05%	0.34%	0.46%	95.64%	99.75%
FREQ 15	0.20	0.78	13.97%	15.10%	2.03%	2.86%	0.41%	0.46%	95.65%	99.75%
FREQ 16	0.21	0.82	14.05%	14.93%	2.02%	2.88%	0.37%	0.43%	95.65%	99.75%
FREQ 17	0.27	0.97	14.02%	14.41%	1.84%	2.92%	0.31%	0.61%	95.64%	99.75%
FREQ 18	0.27	0.94	13.85%	13.86%	1.75%	3.56%	0.38%	0.52%	95.65%	99.75%
FREQ 19	0.30	0.90	13.20%	14.21%	1.79%	3.54%	0.31%	0.54%	95.65%	99.75%
FREQ 20	0.27	0.75	13.26%	15.16%	1.74%	3.26%	0.43%	0.33%	95.65%	99.75%
FREQ 21	0.34	0.73	13.38%	15.47%	1.71%	3.14%	0.19%	0.54%	95.64%	99.75%
FREQ 22	0.33	0.73	13.93%	14.90%	1.62%	3.33%	0.24%	0.57%	95.65%	99.75%
FREQ 23	0.25	0.68	13.34%	15.09%	1.98%	3.34%	0.31%	0.52%	95.65%	99.75%
FREQ 24	0.30	0.75	13.71%	14.62%	1.81%	3.40%	0.34%	0.59%	95.65%	99.75%
FREQ 25	0.43	0.93	13.11%	14.77%	1.78%	3.39%	0.08%	0.77%	95.65%	99.75%
FREQ 26	0.42	0.94	12.78%	14.70%	1.74%	3.32%	0.14%	0.80%	95.65%	99.75%
FREQ 27	0.47	1.04	12.29%	14.51%	1.76%	3.30%	0.15%	0.83%	95.65%	99.75%
OVERALL	0.29	0.68	14.14%	15.00%	1.80%	3.04%	0.20%	0.51%	95.64%	99.75%

Table IV- Results of a leave-one-out cross-validation of a LORETA normative database (from Thatcher et al, 2005b)

The results of a leave-one-out cross-validation are published in Thatcher et al (2005a; 2005b).

Another method of establishing content and construct validity of a LORETA normative database is to test the accuracy of the database using patients with confirmed pathologies where the location of the pathology is known by other imaging methods, e.g., CT-scan or MRI or PET, etc. Validity is estimated by the extent that there is a high correspondence between the location of the confirmed pathology and the location of the 3-dimensional sources of the EEG that correspond to the location of the pathology. Here is a partial list of studies showing concordance validity with fMRI and LORETA (Mobascher et al, 2009a; 2009b; Esposito et al, 2009a; 2009b; Brookings et al, 2009; Yoshioka et al, 2008; Schulz et al, 2008) and between PET and LORETA (Horacek et al, 2007; Hu et al, 2007; Zumsteg et al, 2005; Tišlerová et al, 2005; Kopeček et al, 2005; Pizzagalli et al, 2004) and between SPECT and LORETA (Korn et al, 2005). Figure 12 shows an example of the EEG from an epilepsy patient in which maximal epileptic discharges are present in the left temporal, left parietal and left occipital regions. Content

validity of LORETA is established by the fact that the maximum amplitude of epilepsy was in the left temporal lobe lead (T5) at 3 Hz as measured by the FFT and the Z scores from the scalp surface. The sources were localized to Brodmann area 22 left superior temporal gyrus and Brodmann area 13 of the left insular cortex.



LORETA is low resolution electromagnetic tomography (est. 2 – 4 cm resolution) and precise millimeter localization of epileptic foci is beyond the resolution of LORETA. Nonetheless, verification of the surface EEG with 3-dimensional source currents illustrates the use of hypotheses as to the expected hemisphere and regions based on the

surface EEG. In this case the hypothesis from the surface EEG was that there is an expected source in the left temporal regions (Brodmann areas were predicted beforehand) and this hypothesis was confirmed. This is an example of specificity of a Z score normative database in which 3-dimensional hypotheses are formed (and thus planned comparisons) based on the surface EEG and the hypothesis is then tested using LORETA.

Figure 13 (Top) shows an example of the EEG from a TBI patient with a right hemisphere hematoma. The maximum amplitude of slow waves (1- 6 Hz) was in the right pre-frontal (C4), right parietal (P4) as well as right occipital regions (O2) as measured by the FFT and the Z scores from the scalp surface.

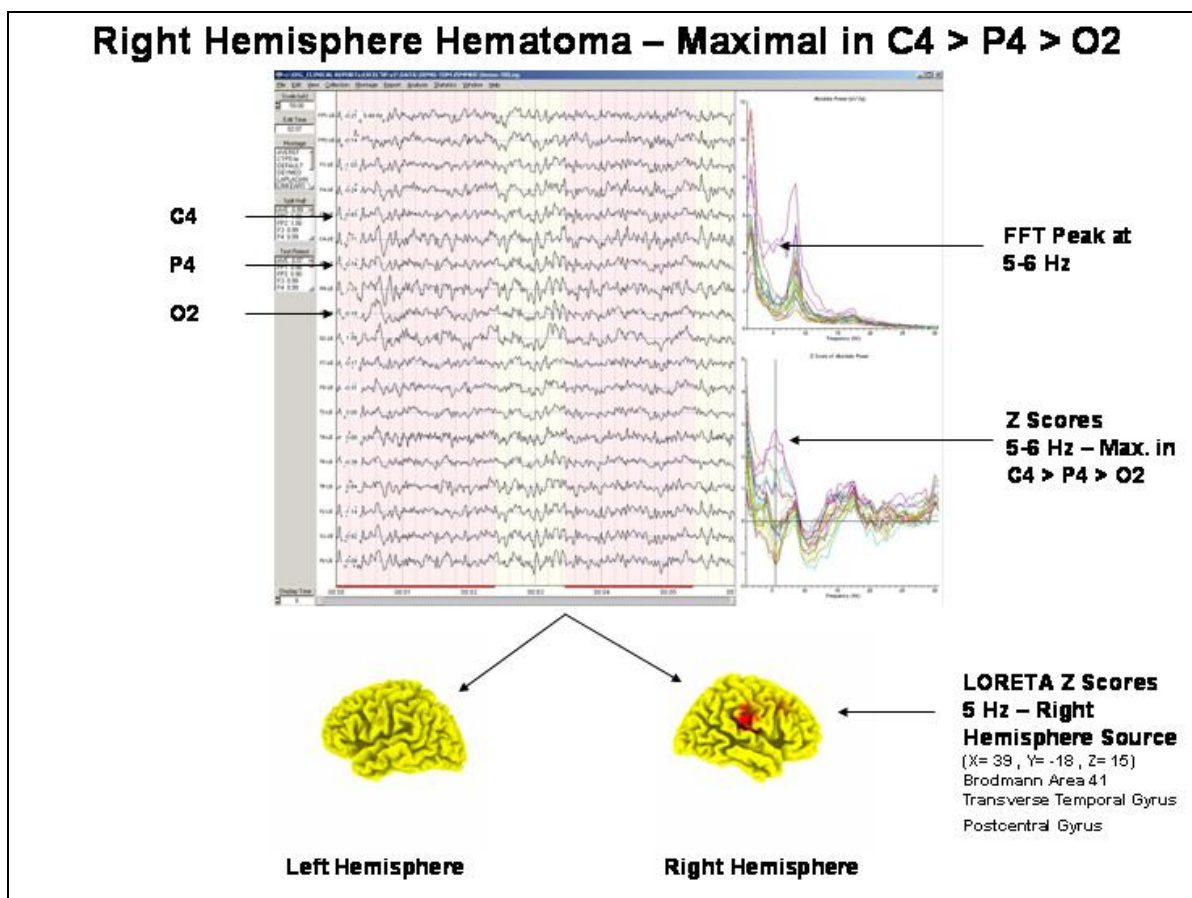
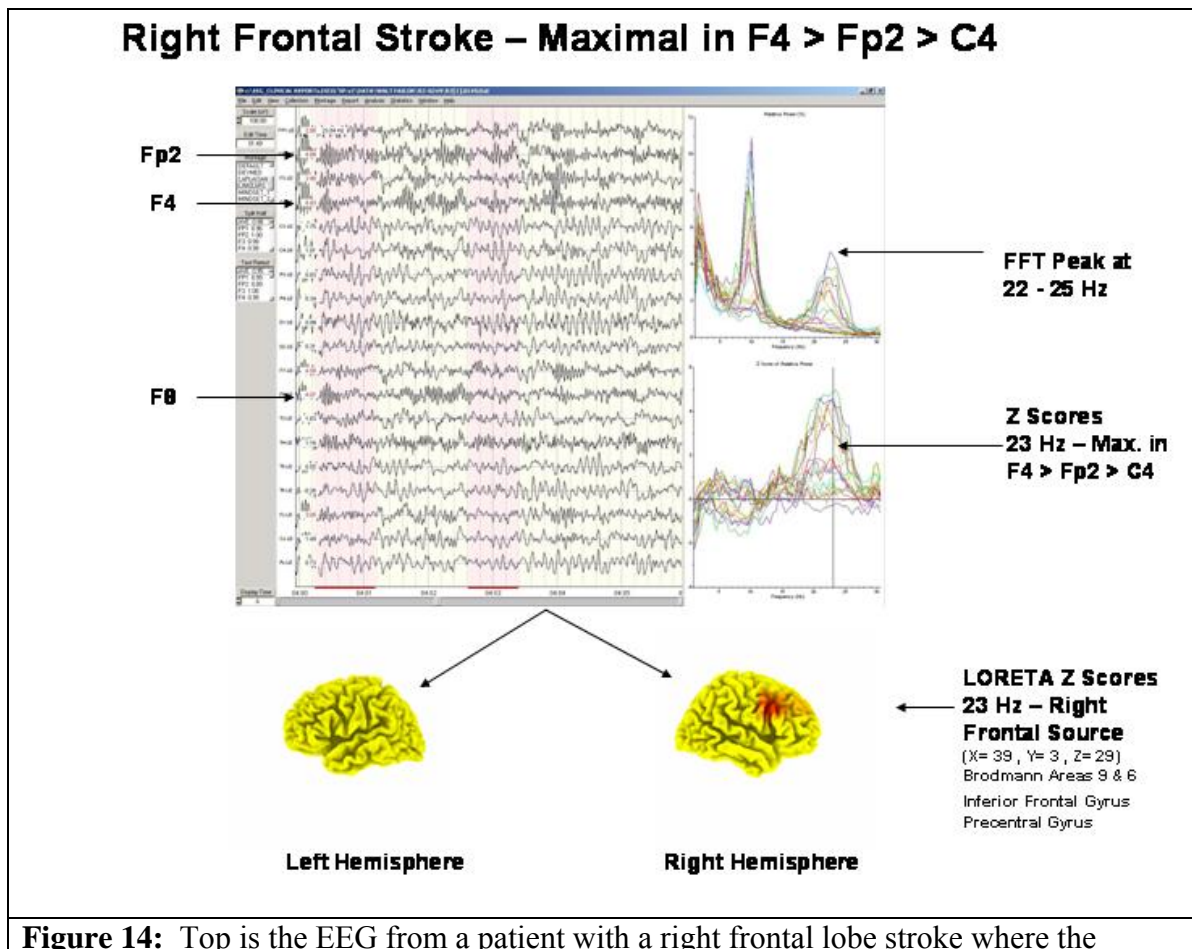


Figure 13: Top is the EEG from a patient with a right hemisphere hematoma where the maximum slow waves are present in C4, P4 and O2. The FFT power spectrum from 1 to 30 Hz and the corresponding Z scores of the surface EEG are shown in the right side of the EEG display. Bottom, are the left and right hemisphere displays of the maximal Z scores using LORETA. It can be seen that only the right hemisphere has statistically significant Z values. Planned comparisons and hypothesis testing based on the frequency and location of maximal deviation from normal on the surface EEG are confirmed by the LORETA Z score normative analysis. (from Thatcher et al, 2005b)

Figure 13 (Bottom) shows the Z scores in LORETA slices in the right hemisphere hematoma patient which were consistent with the surface EEG deviation from normal by being in the right hemisphere and near to the area of maximal damage. The maximum Z scores were present in the right post-central gyrus at 5 Hz and were localized to Brodmann area 43 right post-central gyrus as well as Brodmann areas 13 right insula cortex and 41 right transverse temporal gyrus.

Figure 14 (Top) shows an example of the EEG from a right hemisphere stroke patient. The maximum Z scores from the scalp EEG were in the right anterior frontal regions (F4 & Fp2) at 23 Hz. It can be seen that the maximum Z scores were present in the right frontal regions at 23 Hz and the Key Institute Talairach Atlas were maximally localized to Brodmann area 9 right inferior frontal gyrus as well as Brodmann area 6 right frontal pre-central gyrus. This is another example of validation of a LORETA Z score normative database in which 3-dimensional hypotheses are formed (and thus planned comparisons) based on the surface EEG and the hypothesis is then tested using LORETA.



maximum slow waves are present in F4 and Fp2. The FFT power spectrum from 1 to 30 Hz and the corresponding Z scores of the surface EEG are shown in the right side of the EEG display. Bottom, are the left and right hemisphere displays of the maximal Z scores using LORETA. It can be seen that only the right hemisphere has statistically significant Z values. Planned comparisons and hypothesis testing based on the frequency and location of maximal deviation from normal on the surface EEG are confirmed by the LORETA Z score normative analysis. (from Thatcher et al, 2005b)

Construct Validity of a LORETA normative database based on the smoothness at 1 Hz Resolution and Regions of Interest (ROIs)

A smooth distribution of Z scores with maxima near to the location of the confirmed injury is expected if parametric statistics using LORETA are valid. This is an example of construct validity. Figure 15 is a graph of the rank order of Z scores for different 1 Hz frequency bands from 1 to 10 Hz for the 2,394 current source values in the right hemisphere hematoma patient in figure 15. It can be seen that the rank ordering of the Z scores is smooth and well-behaved at each 1 Hz frequency analysis with maximum Z score deviation at 2 – 6 Hz which is the same frequency band in which the surface EEG was most deviant from normal (see Figure 13). A smooth rank ordering of Z scores is expected if parametric statistical analysis is valid.

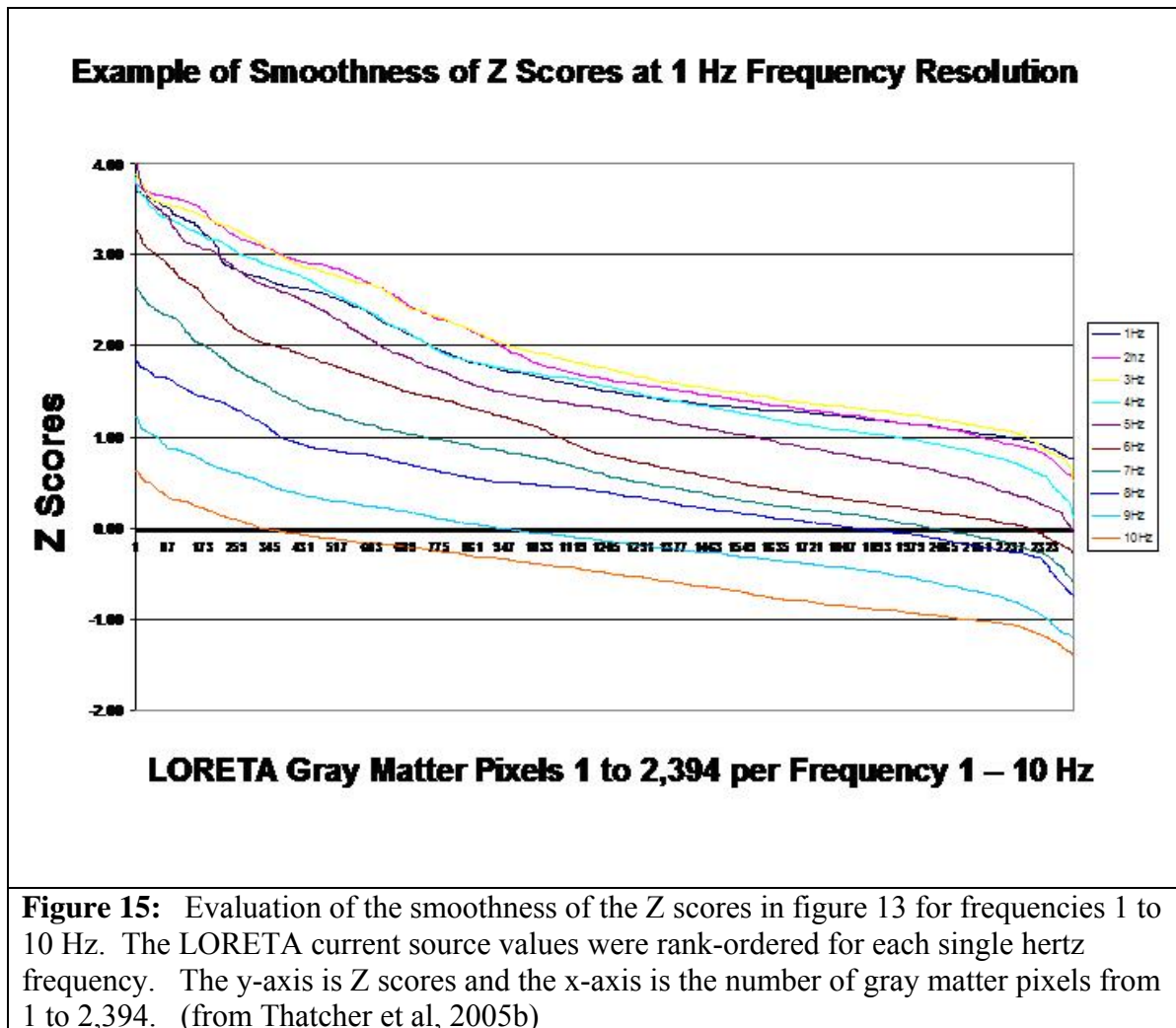


Figure 15: Evaluation of the smoothness of the Z scores in figure 13 for frequencies 1 to 10 Hz. The LORETA current source values were rank-ordered for each single hertz frequency. The y-axis is Z scores and the x-axis is the number of gray matter pixels from 1 to 2,394. (from Thatcher et al, 2005b)

Reliability Defined

Reliability is the extent to which an experiment, test, or any measuring procedure yields the same result on repeated trials. Researchers and clinicians would be unable to satisfactorily draw conclusions, formulate theories, or make claims about the generalizability of their research without the agreement of independent and replicable observations nor to be able to replicate research procedures, or use research tools and procedures that yield consistent measurements. The measurement of any phenomenon always contains a certain amount of chance error. The null hypothesis in any test of reliability is where reliability = 0, that is, repeated measurements of the same phenomenon never duplicate each other and they are not consistent from measurement to measurement. The Type I and Type II errors inherent in the reliability of a sample of

digital EEG and/or qEEG can be measured in different ways. An acceptable level of reliability depends on the intended application of the method and on the tolerance of error.

There are various ways to measure reliability such as: 1- the retest method (stability over time), 2-alternative-form method, 3- internal consistency and 4- split-halves method (Carmines et al, 1979). The particular method of computing reliability depends on the circumstances and/or personal choice. It is possible to have a measure that has high reliability but low validity, that is, one that is consistent in getting wrong information or is consistent in missing the mark. It is also possible for low reliability and low validity, that is, inconsistent and never on target. “Test, re-test reliability” also called “stability reliability” is a commonly used method of reliability testing in qEEG and is generally defined as the agreement of measuring instruments over time. Alternative-form reliability is when different measures provide similar results, for example, EEG coherence and EEG phase lock duration or coherences vs comodulation, etc. To determine stability, a measure or test is repeated on the same subjects at different points in time. Results are compared and correlated with the initial test to give a measure of stability and to detect changes. The test re-test reliability statistic is a good method to detect drowsiness when comparing the beginning of the EEG recording to the end of a lengthy recording with eyes closed. For example, if there is no dramatic change in state between the beginning and end of the recording than one would expect high test re-test reliability (e.g., > 0.9). On the other hand, if a patient is drowsy or sleeping near the end of the recording, then one would expect the test re-test reliability between the beginning of the record to be low (e.g., < 0.9).

Reliability of EEG Autopower Spectrum

The autopower spectrum is the real part of the power spectrum that measures the amount of energy in a complex wave form at each frequency. The units are micovolts squared per cycle per second or $\mu V^2/Hz$. Amplitude or magnitude is simply the square root of power and the same reliability measures are used for both power and amplitude. The scientific literature demonstrating high reliability (e.g., > 0.9) of quantitative EEG is

diverse and quite large and can be read by visiting the National Library of Medicine's database at:

<https://www.ncbi.nlm.nih.gov/sites/entrez?db=pubmed> and use the search terms "EEG and Reliability" and there are a total of 368 citations and a quick review of the abstracts shows that the vast majority if not all of these studies are qEEG studies and demonstrate high test re-test reliability of the qEEG. Below are a small but representative sample of some of the studies demonstrating high reliability with sample lengths as short as 20 seconds (Arruda et al 1996; Burgess A, and Gruzelier, 1993; Corsi-Cabrera et al, 1997; Gasser et al, 1985; 1988a; Hamilton-Bruce et al, 1991; Chabot et al, 1996; Pollock et al, 1991; Fernández et al, 1993; John et al, 1987; 1988; Harmony et al, 1993; Lund et al, 1995; Duffy et al, 1994; Salinsky et al, 1991; McEvoy et al, 2000; Näpflin et al, 2007; 2008; Towers and Allen, 2009; Van Albada et al, 2007).

Gasser et al (1985, pg. 312) concluded: "20 sec of activity are sufficient to reduce adequately the variability inherent in the EEG"

Salinski et al (1991, pg. 382) concluded: "Correlation coefficients for broad band features averaged 0.92 over the 5 min retest interval and 0.84 over the 12-16 and "Coefficients based on 60 sec records were marginally higher than those of 40 or 20 sec records."

Corsi-Cabrera et al (1997, pg. 382) concluded: "The within-subject stability was assessed calculating multiple correlation coefficients between all EEG features of the eleven sessions of each subject: R-values ranged from 0.85 to 0.97."

Pollock et al (1991, pg. 20) concluded: "the generally higher reliabilities of absolute, as opposed to relative, amplitude measures render them preferable in clinical research."

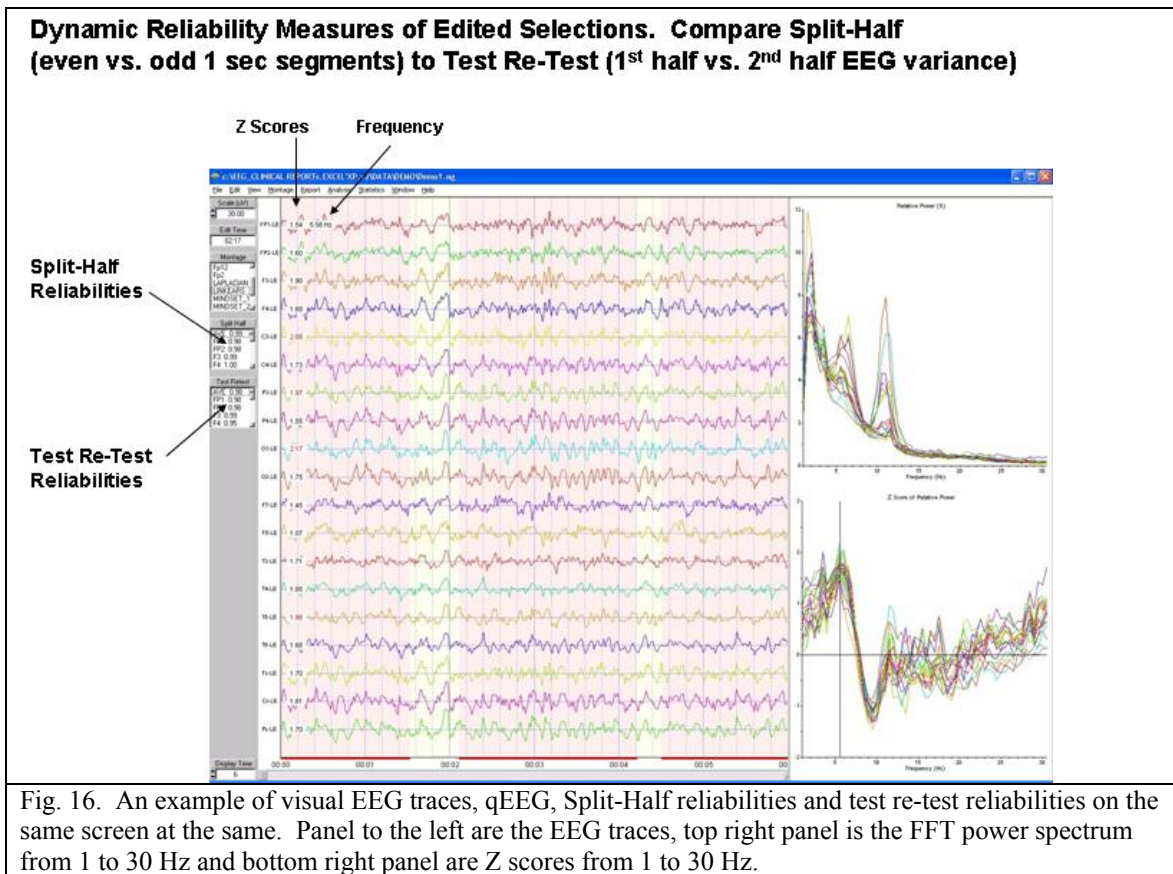
EEG spectral stability over a one year period was recently studied by Näpflin and colleagues with test re-test reliability > 0.9 and they concluded that qEEG intra-individual reliability is very high:

“Out of all 2400 pairwise comparisons 99.3% were correct, with sensitivity 87.5% and specificity 99.5%. The intra-individual stability is high compared to the inter-individual variation. Thus, interleaved EEG-fMRI measurements are valid. Furthermore, longitudinal effects on cognitive EEG can be judged against the intra-individual variability in subjects.” (Näpflin et al, 2008, pg. 2519).

A recent study by Van Albada and colleagues evaluated the variable contributions of “state” and “trait” by conducting test re-test reliability measures of the qEEG recorded from subject each week for six weeks and some subjects for as long as a year and concluded:

“About 95% of the maximum change in spectral parameters was reached within minutes of recording time, implying that repeat recordings are not necessary to capture the bulk of the variability in EEG spectra.” Van Albada et al (2007, pg. 279).

In general, the test re-test reliability of qEEG is an exponential function of sample length in which 20 second epochs are approximately 0.8 reliable, 40 seconds approx. 0.9 reliable and 60 seconds asymptotes at approx. 0.95 reliability. Figure 16 shows an example of visual EEG traces (non-qEEG) and qEEG (right panels) on the same computer screen at the same time. Reliability measures



of selected segments of the EEG are immediately displayed on the left side of the display. In this way professionals can immediately evaluate the test re-test reliability of their artifact free selections and use the qEEG analyses as a micro-analysis or fine grain analysis of the EEG traces. If test re-test reliabilities are > 0.9 and there is no evidence of drowsiness or artifact in the record then further quantitative analyses can be performed.

Reliability of EEG Coherence

As mentioned previously, coherence is itself a statistical measure of reliability because it is a measure of the stability of phase differences between two EEG time series. If the phase difference is unreliable, i.e., phase differences are randomly changing from time sample to time sample, then coherence = 0. If the phase differences are unchanging then coherence = 1. High test re-test reliability of EEG coherence has been reported over the years when coherence is correctly computed even though more statistical samples are often required in order to obtain statistical sufficiency. If regions of the brain

are weakly coupled or disconnected, then coherence has low values within a subject as well as low test re-test reliability across experiments and across subjects as expected. If regions of the brain are strongly coupled and coherence exhibits statistically significant values then coherence typically also exhibits high test re-test reliability (the greater the coherence then the more within session and between session reliability by definition). Adey et al (1961) were among the first to measure the test re-test reliability of EEG coherence with values > 0.8 . Subsequently, high re-test reliability of EEG coherence (0.8 to 0.95) was reported by John (1977); John et al (1987); Chabot et al (1996); Gasser et al (1988a); Harmony et al, 1993; Thatcher et al, 1986; 2003; and Corsi-Cabrera et al (2007). Gudmondsson et al (2007) reported low test re-test reliability of coherence because of an invalid computation of coherence due to the use of an average reference. If the authors used a common reference and coherence was low, e.g., < 0.2 then this means that two brain regions are reliably disconnected. If the reader finds any study that claims that coherence has low reliability, immediately examine the methods section and see if the authors used an average reference or a Laplacian reference or ICA to create a new time series and if so, then dismiss the study because they used an invalid method of measuring coherence in the first place. Remember, reliability is irrelevant if the measure is not valid to begin with.

Summary

The fact that qEEG meets high standards of reliability and validity is demonstrated by hundreds of paper reviewed journal articles a few of which are cited in this review. The critics of qEEG are those that rely solely on “Eye-Ball” examination of the EEG traces and are biased against and opposed to the use of computers to improve the accuracy, validity and reliability of the electroencephalogram (Nuwer, 1997). The American Academy of Neurology (ANN) position paper (Nuwer, 1997) categorized qEEG as “experimental” for a wide range of clinical disorders because of the blanket assertion that qEEG is “unreliable” without citing a single study to refute the scientific literature that demonstrates high reliability and validity. Hopefully, this review will help those that use qEEG for clinical purposes to refute the false claims of those that make blanket statements that the qEEG is invalid and unreliable by responding with solid

scientific evidence that proves the opposite. It is the responsibility of those that use qEEG technology to respond to false claims by citing facts and citing the scientific literature when ever possible.

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Figure Legends

Figure One - shows an example of four 1 uV and 5 Hz sine waves with the second to the 4th sine wave shifted by 30 degrees. Gaussian noise is added incrementally to channels 2 to 4. Channel 2 = 1 uV signal + 2 uV of noise, channel 3 = 1 uV signal + 4 uV of noise and channel 4 = 1 uV signal + 6 uV of noise. Nineteen channels were used in the analyses of coherence in 2 uV of noise increments. The FFT analysis is the mean of thirty 2 second epochs sampled at 128 Hz.

Figure Two- Top is coherence (y-axis) vs signal-to-noise ratio (x-axis). Bottom is phase angle on the y-axis and signal-to-noise ratio on the x-axis. Phase locking is minimal or absent when coherence is less than approximately 0.2 or 20%. The sample size was 60 seconds of EEG data and smoother curves can be obtained by increasing the epoch length.

Figure Three - Left top is coherence (y-axis) vs signal-to-noise ratio (x-axis) with a 30^o phase shift as shown in figure 2 using the average reference. The left bottom is phase differences in degrees in the y-axis and the x-axis is the signal-to-noise ratio using the average reference. The

right top graph is coherence (y-axis) vs signal-to-noise ratio (x-axis) using the Laplacian montage. The right bottom is phase difference on the y-axis and signal-to-noise on the x-axis using the Laplacian montage. In both instances, coherence drops off rapidly and is invalid with no linear relationship between signal and noise. The bottom graphs show that both the average reference and the Laplacian montage fails to track the 30° phase shift that was present in the original time series. In fact, the phase difference is totally absent and unrepresented when using an average reference or a Laplacian montage and these simulations demonstrate that the average reference and the Laplacian montage are not physiologically valid because they do not preserve phase differences or the essential time differences on which the brain operates.

Figure Four- Demonstration of distortions in phase differences in a test using 20 deg increments of phase difference with respect to Fp1. The solid black line is using a Linked Ears common reference which accurately shows the step by step 20 deg. Increments in phase difference. The average reference (dashed blue line) and the Laplacian (dashed red line) significantly distort the phase differences.

Figure Five- Example of Gaussian Cross-Validation of EEG Normative Database (from Thatcher et al, 2003).

Figure Six - Illustration of method of computing error rates or sensitivity of a normative EEG database based on the cross-validation deviation from Gaussian (from Thatcher et al, 2003a).

Figure Seven – Example of predictive and content validity by clinical correlations of qEEG with Neuropsychological test scores (Thatcher et al, 2001).

Figure Eight - Example of content validity demonstrated by statistically significant correlations between full scale I.Q. and qEEG (from Thatcher et al, 2005c).

Figure Nine - Example of “Planned comparisons” using hypothesis creation prior to launching LORETA. Content and construct validity are present because the patient was hit on the right parietal lobe and the right parietal lobe shows deviant EEG activity (e.g., > 2 st. dev.) Further construct validity is established by LORETA analyses that confirm anatomical hypotheses based on the surface EEG locations and frequencies of deviance.

Figure Ten - An example of construct validity of the qEEG to correlate with the MRI in the estimate of traumatic brain injury (adapted from Thatcher et al, 1998a; 1998b).

Figure Eleven- The distribution of the Z scores of the current source density LORETA values at 1 Hz resolution. The y-axis is the number or count and the x-axis is the Z Score, defined as the

mean – each value in each of the 2,394 pixels divided by the standard deviation (from Thatcher et al, 2005b).

Figure Twelve - Top is the EEG from a patient with Left Temporal Lobe epilepsy where the maximum spike and waves are present in T5, O1, P3 and T3. The FFT power spectrum and the corresponding surface EEG Z scores are shown in the top right side. Bottom, are the left and right hemisphere displays of the maximal Z scores using LORETA. It can be seen that only the left temporal lobe has statistically significant Z values. Planned comparisons and hypothesis testing based on the frequency and location of maximal deviation from normal on the surface EEG are confirmed by the LORETA Z score normative analysis (from Thatcher et al, 2005b).

Figure Thirteen - Top is the EEG from a patient with a right hemisphere hematoma where the maximum slows waves are present in C4, P4 and O2. The FFT power spectrum from 1 to 30 Hz and the corresponding Z scores of the surface EEG are shown in the right side of the EEG display. Bottom, are the left and right hemisphere displays of the maximal Z scores using LORETA. It can be seen that only the right hemisphere has statistically significant Z values. Planned comparisons and hypothesis testing based on the frequency and location of maximal deviation from normal on the surface EEG are confirmed by the LORETA Z score normative analysis. (from Thatcher et al, 2005b).

Figure Fourteen - Top is the EEG from a patient with a right frontal lobe stroke where the maximum slows waves are present in F4 and Fp2. The FFT power spectrum from 1 to 30 Hz and the corresponding Z scores of the surface EEG are shown in the right side of the EEG display. Bottom, are the left and right hemisphere displays of the maximal Z scores using LORETA. It can be seen that only the right hemisphere has statistically significant Z values. Planned comparisons and hypothesis testing based on the frequency and location of maximal deviation from normal on the surface EEG are confirmed by the LORETA Z score normative analysis. (from Thatcher et al, 2005b).

Figure Fifteen - Figure 15: Evaluation of the smoothness of the Z scores in figure 13 for frequencies 1 to 10 Hz. The LORETA current source values were rank-ordered for each single hertz frequency. The y-axis is Z scores and the x-axis is the number of gray matter pixels from 1 to 2,394. (from Thatcher et al, 2005b)

Figure Sixteen - An example of visual EEG traces, qEEG, Split-Half reliabilities and test re-test reliabilities on the same screen at the same. Panel to the left are the EEG traces, top right panel is the FFT power spectrum from 1 to 30 Hz and bottom right panel are Z scores from 1 to 30 Hz.

7.0 – Table Legends

Table I - Cross Validation of EEG Normative Database (from Thatcher et al, 2003).

Table II - Normative EEG database sensitivities for different age groups at +/- 2 standard deviations and +/- 3 standard deviations (from Thatcher et al, 2003a).

Table III - Examples of predictive validity by clinical correlations between qEEG and intelligence (WISC-R) and academic achievement tests (WRAT) (from Thatcher et al, 2003a).

Table IV - Results of a leave-one-out cross-validation of a LORETA normative database (from Thatcher et al, 2005b)