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AN OVERVIEW ON ASSESSING AGREEMENT WITH CONTINUOUS MEASUREMENTS

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Reliable and accurate measurements serve as the basis for evaluation in many scientific disciplines. Issues related to reliable and accurate measurement have evolved over many decades, dating back to the nineteenth century and the pioneering work of Galton (1886), Pearson (1896, 1899, 1901), and Fisher (1925). Requiring a new measurement to be identical to the truth is often impractical, either because (1) we are willing to accept a measurement up to some tolerable (or acceptable) error, or (2) the truth is simply not available to us, either because it is not measurable or is only measurable with some degree of error. To deal with issues related to both (1) and (2), a number of concepts, methods, and theories have been developed in various disciplines. Some of these concepts have been used across disciplines, while others have been limited to a particular field but may have potential uses in other disciplines. In this paper, we elucidate and contrast fundamental concepts employed in different disciplines and unite these concepts into one common theme: assessing closeness (agreement) of observations. We focus on assessing agreement with continuous measurements and classify different statistical approaches as (1) descriptive tools; (2) unscaled summary indices based on absolute differences of measurements; and (3) scaled summary indices attaining values between -1 and 1 for various data structures, and for cases with and without a reference. We also identify gaps that require further research and discuss future directions in assessing agreement.

Key Words: Accuracy; Agreement; Coefficient of individual agreement; Concordance correlation coefficient; Coverage probability; Generalizability; Intraclass correlation coefficient; Limits of agreement; Method comparison; Precision; Reliability; Repeatability; Reproducibility; Tolerance interval; Total deviation index; Validity.

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1. INTRODUCTION

In social, behavioral, physical, biological, and medical sciences, reliable and accurate measurements serve as the basis for evaluation. As new concepts, theories, and technologies continue to develop, new scales, methods, tests, assays, devices, and instruments for evaluation become available for measurement. Because errors are inherent in every measurement procedure, one must ensure that the measurement is reliable and accurate before it is used in practice. The issues related to reliable and accurate measurement have evolved over many decades, dating back to the nineteenth century and the pioneering work of Galton (1886), Pearson (1896, 1899, 1901), and Fisher (1925): from the intraclass correlation coefficient (ICC) that measures reliability (Bartko, 1966; Fisher, 1925; Galton, 1886; Pearson, 1896; Shrout and Fleiss, 1979; Vangeneugden et al., 2004) and the design of reliability studies (Donner, 1998; Dunn, 2002; Shoukri et al., 2004), to generalizability extending the concept of ICC (Brennan, 2001; Cronback, 1951; Cronbach et al., 1972; Lord and Novick, 1968; Vangeneugden et al., 2005); from the International Organization for Standardizations (ISO) (1994) guiding principle on accuracy of measurement (ISO 5725-1) to the Food and Drug Administrations (FDA) (2001) guidelines on bioanalytical method validation; and including various indices to assess the closeness (agreement) of observations (Bland and Altman, 1986, 1995, 1999; Barnhart et al., 2002, 2005a; Carrasco and Jover, 2003a; Choudhary and Nagaraja, 2004; Dunn, 2004; Haber and Barnhart, 2006; King and Chinchilli, 2001a; Lin, 1989, 2000a, 2003; Lin et al., 2002; Shrout, 1998).

In the simplest intuitive terms, reliable and accurate measurement may simply mean that the new measurement is the same as the truth or agrees with the truth. However, requiring the new measurement to be identical to the truth is often impractical, either because (1) we are willing to accept a measurement up to some tolerable (or acceptable) error or (2) the truth is simply not available to us (either because it is not measurable or because it is only measurable with some degree of error). To deal with issues related to both (1) and (2), a number of concepts, methods, and theories have been developed in different disciplines. For continuous measurement, the related concepts are accuracy, precision, repeatability, reproducibility, validity, reliability, generalizability, agreement, etc. Some of these concepts (e.g., reliability) have been used across different disciplines. However, other concepts, such as generalizability and agreement, have been limited to a particular field but may have potential uses in other disciplines.

In this paper, we describe and contrast the fundamental concepts used in different disciplines and unite these concepts into one common theme: assessing closeness (agreement) of observations. We focus on continuous measurements and summarize methodological approaches for expressing these concepts and methods mathematically, and discuss the data structures for which they are to be used, both for cases with and without a reference (or truth). Existing approaches for expressing agreement are organized in terms of the following: (1) descriptive tools, such as pairwise plots with a 45-degree line and Bland and Altman plots (Bland and Altman, 1986); (2) unscaled summary indices based on absolute differences of measurements, such as mean squared deviation including repeatability coefficient and reproducibility coefficient, limits of agreement (Bland and Altman, 1999), coverage probability, and total deviation index (Lin et al., 2002); and (3) scaled summary

indices attaining values between -1 and 1 , such as the intraclass correlation coefficient, concordance correlation coefficient, coefficient of individual agreement, and dependability coefficient.

These approaches were developed for one or more types of the following data structure: (1) two or more observers without replications; (2) two or more observers with replications; (3) one or more observer is treated as a random or fixed reference; (4) longitudinal data where observers take measurements over time; (5) data where covariates are available for assessing the impact of various factors on agreement measures. We discuss the interpretation of the magnitude of the agreement values on using the measurements in clinical practice. We also identify gaps that require further research, and discuss future directions in assessing agreement. In Section 2, we present definitions of different concepts used in the literature and provide our critique. Statistical approaches are presented in Section 3 where various concepts are used. We conclude with a summary and discussions of future directions in Section 4.

2. CONCEPTS

2.1. Accuracy and Precision

In Merriam Webster's dictionary, *accuracy* and *precision* are synonyms. *Accuracy* is defined as "freedom from mistake or error" or "conformity to truth or to a standard" or "degree of conformity of a measure to a standard or a true value." *Precision* is defined as "the quality of being exactly or sharply defined" or "the degree of refinement with which a measurement is stated." The "degree of conformity" and "degree of refinement" may mean the same thing. The subtle difference between these two terms may lie in whether a truth or a reference standard is required or not.

2.1.1. Accuracy. Historically, accuracy has been used to measure systematic bias while precision has been used to measure random error around the expected value. Confusion regarding the use of these two terms continues today because of the existence of different definitions and because of the fact that these two terms are sometimes used interchangeably. For example, the US Food and Drug Administration (FDA) guidelines on bioanalytical method validation (1999) defined *accuracy* as the closeness of mean test results obtained by the method to the true value (concentration) of the analyte. The deviation of the mean from the true value, i.e., systematic bias, serves as the measure of accuracy. However, in 1994, the International Organization for Standardization (ISO) used accuracy to measure both systematic bias (trueness) and random error. In ISO 5725 (1994), the general term *accuracy* was used to refer to both trueness and precision, where "trueness" refers to the closeness of agreement between the arithmetic mean of a large number of test results and the true or accepted reference value, and "precision" refers to the closeness of agreement between test results. In other words, accuracy involves both systematic bias and random error, because "trueness" measures systematic bias. The ISO 5725 (1994) acknowledged that:

"The term accuracy was at one time used to cover only the one component now named trueness, but it became clear that to many persons it should imply the total

displacement of a result from a reference value, due to random as well as systematic effects. The term bias has been in use for statistical matters for a very long time, but because it caused certain philosophical objections among members of some professions (such as medical and legal practitioners), the positive aspect has been emphasized by the invention of the term “trueness”.

Despite the ISO's effort to use one term (accuracy) to measure both systematic and random errors, the use of accuracy for measuring the systematic bias, and precision for measuring random error, is commonly encountered in the literature of medical and statistical research. For this reason, we will use *accuracy* to stand for systematic bias in this paper, where one has a “true sense of accuracy” (systematic shift from truth) if there is a reference, and a “loose sense of accuracy” (systematic shift from each other) if no reference is used for comparison. Thus, the “true sense of accuracy” used in this paper corresponds to the FDA's accuracy definition and the ISO's trueness definition. Ideally and intuitively, the accepted reference value should be the true value, because one can imagine that the true value has always existed, and the true value should be used to judge whether there is an error. However, in social and behavioral sciences, the true value may be an abstract concept, such as intelligence, which may only exist in theory and may thus not be amenable to direct measurement. In biomedical sciences, the true value may be measured with a so-called gold standard that may also contain small amount of systematic and/or random error. Therefore, it is very important to report the accepted reference, whether it is the truth or subject to error (including the degree of systematic and random error if known). In this paper, we only consider the case where the reference or gold standard is measured with error.

2.1.2. Precision. The FDA (1999) defined *precision* as the closeness of agreement (degree of scatter) between a series of measurements obtained from multiple sampling of the same homogeneous sample under the prescribed conditions. Precision is further subdivided into within-run, intra-batch precision or repeatability (which assesses precision during a single analytical run) and between-run, inter-batch precision or repeatability (which measures precision over time, and may involve different analysts, equipment, reagents, and laboratories).

ISO 5725 (1994) defined *precision* as the closeness of agreement between independent test results obtained under stipulated conditions. ISO defined repeatability and reproducibility as precision under the repeatability and reproducibility conditions, respectively (see Section 2.2).

The key phrase is “under the prescribed conditions” or “under stipulated conditions.” It is therefore important to emphasize the conditions used when reporting precision. Precisions are only comparable under the same conditions.

2.2. Repeatability and Reproducibility

Repeatability and reproducibility are two special kinds of precision under two extreme conditions and they should not be used interchangeably. As defined below, repeatability assesses pure random error due to “true” replications and reproducibility assesses closeness between observations made under condition other than pure replication, e.g., by different labs or observers. If precision is expressed

by imprecision such as standard deviation, repeatability is always smaller than or equal to reproducibility (see below for definition).

2.2.1. Repeatability. The FDA (2001) used the term *repeatability* for both intra-batch precision and inter-batch precision. The ISO defined *repeatability* as the closeness of agreement between independent test results under repeatability conditions that are as constant as possible, where independent test results are obtained with the same methods, on identical test items, in the same laboratory, performed by the same operator, using the same equipment, within short intervals of time.

We use the ISO's definition of repeatability in this paper. To define the term more broadly, *repeatability* is the closeness of agreement between measures under the same condition, where "same condition" means that nothing changed other than the times of the measurements. The measurements taken under the same condition can be viewed as true replicates.

Sometimes the subject does not change over time, as in the case of x-ray slides or blood samples. However, in practice, it may be difficult to maintain the same condition over time when measurements are taken. This is especially true in the social and behavioral sciences, where characteristics or constructs change over time due to learning effects. It is important to ensure that human observers are blinded to earlier measurements of the same quantity. We frequently rely on *believable assumptions* that the same condition is maintained over a short period of time when measurements are taken. It is essential to state *what assumptions* are used when reporting repeatability. For example, when an observer uses an instrument to measure a subject's blood pressure, the same condition means the same observer using the same instrument to measure the same subject's blood pressure, where the subject's blood pressure did not change over the course of multiple measurements. It is unlikely that the subject's blood pressure remains constant over time; however, it is believable that the true blood pressure did not change over a short period time, e.g., a few seconds. Therefore, blood pressures taken in successive seconds by the same observers, using the same instrument on the same subject, may be considered true replicates.

It is important to report repeatability when assessing measurement, because it measures the purest random error that is not influenced by any other factors. If true replicates cannot be obtained, then we have a loose sense of repeatability based on assumptions.

2.2.2. Reproducibility. In 2001, FDA guidelines defined *reproducibility* as the precision between two laboratories. Reproducibility also represents the precision of the method under the same operating conditions over a short period of time. In 1994, the ISO defined *reproducibility* as the closeness of agreement between independent test results under reproducibility conditions under which results are obtained with the same method on identical test items, but in different laboratories with different operators and using different equipment.

We use the ISO's definition of reproducibility in this paper. To define the term more broadly, *reproducibility* is the closeness of agreement between measures under all possible conditions on identical subjects for which measurements are taken.

All possible conditions means any conceivable situation for which a measurement will be taken in practice, including different laboratories, different observers, etc. However, if multiple measurements on the same subject cannot be taken at the same time, one must ensure that the thing being measured (e.g. a subject's blood pressure) does not change over time when measurements are taken in order to assess reproducibility.

2.3. Validity and Reliability

The concepts of accuracy and precision originated in the physical sciences, where direct measurements are possible. The similar concepts of validity and reliability are used in the social sciences, where a reference is required for validity but not necessarily required for reliability. As elaborated below, validity is similar to true sense of agreement with both good true sense of accuracy and precision. Reliability is similar to loose sense of agreement with both good loose sense of accuracy and precision. Historically, validity and reliability have been assessed via scaled indices.

2.3.1. Validity. In social, educational, and psychological testing, *validity* refers to the degree to which evidence and theory support the interpretation of measurement (AERA et al., 1999). Depending on the selection of the accepted reference (criterion or gold standard), there are several types of validity such as *content*, *construct*, *criterion* validity (AERA et al., 1999; Goodwin, 1997; Hand, 2004; Kraemer et al., 2002; Molenberghs et al., 2007). *Content* validity is defined as the extent to which the measurement method assesses all the important content. *Face* validity is similar to *content* validity, and is defined as the extent to which the measurement method assesses the desired content at face. Face validity may be determined by the judgment of experts in the field. *Construct* validity is used when attempting to measure a hypothetical construct that may not be readily observed, such as anxiety. *Convergent* and *discriminant* validity may be used to assess construct validity by showing that the new measurement is correlated with other measurements of the same construct and that the proposed measurement is not correlated with the unrelated construct, respectively. *Criterion* validity is further divided into *concurrent* and *predictive* validity, where criterion validity deals with correlation of the new measurement with a criterion measurement (such as a gold standard) and predictive validity deals with the correlation of the new measurement with a future criterion, such as a clinical endpoint.

Validity is historically assessed by the correlation coefficient between the new measure and the reference (or construct). If there is no systematic shift of the new measurement from the reference or construct, this correlation may be expressed as the proportion of the observed variance that reflects variance in the construct that the instrument or method was intended to measure (Kraemer et al., 2002). For validation of bioanalytical methods, the FDA (2001) provided guidelines on full validation that involve parameters such as (1) accuracy, (2) precision, (3) selectivity, (4) sensitivity, (5) reproducibility, and (6) stability, when a reference is available. The parameters related to selectivity, sensitivity, and stability may only be applicable in bioanalytical method. When the type of validity is concerned with the closeness (agreement) of the new measurement and the reference, we believe

that an agreement index is better suited than the correlation coefficient for assessing validity. Therefore, a statistical approach for assessing agreement for the case with a reference in Section 3 can be used for assessing validity. Other validity measures that are based on a specific theoretical framework and are not concerned with closeness of observations will not be discussed in Section 3.

2.3.2. Reliability. The concept of reliability has evolved over several decades. It was initially developed in social, behavioral, educational, and psychological disciplines, and was later widely used in the physical, biological, and medical sciences (Bartko, 1966; Donner, 1998; Fisher, 1925; Lord and Novick, 1968; Müller and Büttner, 1994; McGraw and Wong, 1996; Shrout and Fleiss, 1979; Shrout, 1998; Shoukri et al., 2004; Vangeneugden et al., 2004). Rather than reviewing the entire body of literature, we provide our point of view on its development. *Reliability* was originally defined as the ratio of true score variance to the observed total score variance in classical test theory (Cronbach et al., 1972; Lord and Novick, 1968), and is interpreted as the percent of observed variance explained by the true score variance. It was initially intended to assess the measurement error if an observer takes a measurement repeatedly on the same subject under identical conditions, or to measure the consistency of two readings obtained by two different instruments on the same subject under identical conditions. If the true score is the construct, then reliability is similar to the criterion validity. In practice, the true score is usually not available, and in this case, reliability represents the scaled precision. Reliability is often defined with additional assumptions. The following three assumptions are inherently used and are not usually stated when reporting reliability.

- (a) The true score exists but is not directly measurable.
- (b) The measurement is the sum of the true score and a random error, where random errors have mean zero and are uncorrelated with each other and with the true score (both within and across subjects).
- (c) Any two measurements for the same subject are *parallel* measurements.

In this context, *parallel* measurements are any two measurements for the same subject that have the same means and variances. With assumptions (a) and (b), reliability, defined above as the ratio of variances, is equivalent to the *square of the correlation coefficient* between the observed reading and the true score. With assumptions (a) through (c), reliability, as defined above, is equivalent to the correlation of any two measurements on the same subject. This correlation is called *intraclass correlation* (ICC) and was originally defined by Galton (1889) as the ratio of fraternal regression, a correlation between measurements from the same class (in this case, brothers) in study of fraternal resemblance in genetics. Estimations of this ICC based on sample moment (Pearson, 1896, 1899, 1901) and on variance components (Fisher, 1925) were later proposed. Parallel readings are considered to come from the same class and can be represented by a one-way ANOVA model (see Section 3). Reliability expressed in terms of ICC is the most common parameter used across different disciplines. Different versions of ICC used for assessing reliability have been advocated (Bartko, 1966; Eliasziw et al., 1994; Müller and Büttner, 1994; McGraw and Wong, 1996; Shrout and Fleiss, 1979) when

different ANOVA models are used in place of assumptions (b) and (c). We discuss these versions in Section 3.

2.4. Dependability and Generalizability

The recognition that assumptions (b) and (c) in classical test theory are too simplistic prompted the development of generalizability theory (GT) (Cronbach et al., 1972; Shavelson and Webb, 1981, 1991, 1992; Shavelson et al., 1989; Brennan 1992, 2000, 2001). GT is widely known and is used in educational and psychological testing literature; however, it is barely used in medical research despite many efforts to encourage broader use since its introduction by Cronbach et al. (1972). This may be due to the overwhelming statistical concepts involved in the theory and the limited number of statisticians who have worked in this area. Recently, Vangeneugden et al. (2005) and Molenberghs et al. (2007) presented linear mixed model approaches to estimating reliability and generalizability in the setting of a clinical trial.

GT extends classical test theory by decomposing the error term into multiple sources of measurement errors, thus relaxing the assumption of parallel readings. The concept of reliability is then extended to the general concept of generalizability or dependability within the context of GT. In general, two studies (G-study and D-study) are involved, with the G-study aimed at estimating the magnitudes of variances due to multiple sources of variability through an ANOVA model, and the D-study, which uses some or all of the sources of variability from the G-study to define specific coefficients that generalize the reliability coefficient, depending on the intended decisions. In order to specify a G-study, the researcher must define the universe of generalizability *a priori*. The universe of generalizability contains factors with several levels/conditions (finite or infinite) so that researchers can establish the interchangeability of these levels. For example, suppose there are J observers and a researcher wants to know whether the J observers are interchangeable in terms of using a measurement scale on a subject. The universe of generalizability would include the observer as a factor with J levels. This example corresponds to the single-facet design. The question of reliability among the J observers thus becomes the question of generalizability or dependability of the J observers.

To define the generalizability coefficient or dependability coefficient, one must specify a D-study and the type of decision. The *generalizability coefficient* involves the decision based on relative error; i.e., how subjects are ranked according to J observers, regardless of the observed score. The *dependability coefficient* involves the decision based on absolute error; i.e., how the observed measurement differs from the true score of the subject. To specify a D study, the researcher must also decide how to use the measurement scale; e.g., does the researcher want to use the single measurement taken by one of J observers, or the average measurement taken by all J observers? Different decisions will result in different coefficients of generalizability and dependability.

2.5. Agreement

Agreement measures the “closeness” between readings. Therefore, agreement is a **broader** term that contains both accuracy and precision. If one of the readings

is treated as the accepted reference, the agreement is concerning validity. If all of the readings can be assumed to come from the same underlying distribution, then agreement is assessing precision around the mean of the readings. When there is a disagreement, one must know whether the source of disagreement arose from systematic shift (bias) or random error. This is important, because a systematic shift (inaccuracy) usually can be fixed with ease through calibration, while a random error (imprecision) is often a more cumbersome exercise of variation reduction.

In absolute terms, readings agree only if they are identical and disagree if they are not identical. However, readings obtained on the same subject or materials under “same condition” or different conditions are not generally identical, due to unavoidable errors in every measurement procedure. Therefore, there is a need to quantify the agreement or “closeness” between readings. Such quantification is best based on the distance between the readings. Therefore, measures of agreement are often defined as functions of the absolute differences between readings. This type of agreement is called *absolute agreement*. Absolute agreement is a special case of the concept of *relational agreement* introduced by Stine (1989). In order to define a coefficient of relational agreement, one must first define a class of transformations that is allowed for agreement. For example, one can decide that observers are in agreement if the scores of two observers differ by a constant; then, the class of transformation consists of all the functions that add the same constant to each measurement (corresponding to additive agreement). Similarly, in the case where the interest is in linear agreement, observers are said to be in agreement if the scores of one observer are a fixed linear function of those of another. In most cases, however, one would not tolerate any systematic differences between observers. Hence, the most common type of agreement is absolute agreement. In this paper, we only discuss indices based on absolute agreement and direct the readers to the literature (Fagot, 1993; Haber and Barnhart, 2006; Stine, 1989; Zegers, 1986) for relational agreement.

Assessing agreement is often used in medical research for method comparisons (Bland and Altman, 1986, 1995, 1999; Dunn, 2004; St. Laurent, 1998), assay validation, and individual bioequivalence (Lin, 1989, 1992, 2000b, 2003; Lin et al., 2002). We note that the concepts of agreement and reliability may appear different. As pointed out by Vangeneugden et al. (2005) and Molenberghs et al. (2007), agreement assesses the degree of closeness between readings within a subject, while reliability assesses the degree of differentiation between subjects; i.e., the ability to tell subjects apart from each other within a population. It is possible that in homogeneous populations, agreement is high but reliability is low, while in heterogeneous populations, agreement may be low but reliability may be high. This is true if unscaled index is used for assessing agreement while the scaled index is used for assessing reliability, because scaled index often depends on between-subject variability (as shown in Section 3.2) and as a result they may appear to assess the degree of differentiation of subjects from a population. When a scaled index is used to assess agreement, the traditional reliability index is a scaled agreement index (see Section 3). As indicated in Section 3.3.2, the concordance correlation coefficient (CCC), a popular index for assessing agreement, is a scaled index for assessing difference between observations. Under comparison of the CCC and the ICC in Section 3.3.2, the CCC reduces to the ICC under the ANOVA models used to define the ICC. Therefore, reliability assessed by ICC is a scaled agreement index under ANOVA assumptions.

In summary, the concepts used to assess reliable and accurate measurements have a common theme: assessing closeness (agreement) between observations. We therefore review the statistical approaches used to assess agreement and relate the approaches to these concepts in the next section.

3. STATISTICAL APPROACHES FOR ASSESSING AGREEMENT

We now summarize the methodological approaches by which the concepts described in Section 2 are expressed mathematically for different types of data structures. Existing approaches for expressing agreement are organized in terms of the following: (1) descriptive tools, (2) unscaled agreement indices, and (3) scaled agreement indices. Data structures include (1) one reading by each of multiple observers; (2) multiple readings by each of multiple observers; and (3) factors or covariates available that may be associated with the degree of agreement.

We are interested in comparisons of measurements or readings by different observers, methods, instruments, laboratories, assays, devices, etc. on the same subject or sample. For simplicity, we use “observers” as a broad term to stand for either observers, methods, instruments, laboratories, assays or devices, etc.. In general, we treat observers as fixed unless they are indicated as random. We consider the situation that the intended use of the measurement in practice is a single observation made by an observer. Thus, the agreement indices reviewed here are interpreted as measures of agreement between observers to determine whether their single observations can be used interchangeably, although data with multiple observations, such as replications or repeated measures by the same observer, may be used to evaluate the strength of the agreement for single observations.

We consider two distinct situations: (1) the J observers are treated symmetrically where none of them is treated as a reference; (2) one of the observers is a reference. Unless stated otherwise, we can assume that we are discussing issues without a reference observer. If there is a reference observer, we use the J th observer as the reference where the reference is also measured with error.

We use subject to denote subject or sample, where the subjects or samples are randomly sampled from a population. Throughout this section, let Y_{ijk} be the k th reading for subject i made by observer j . In most situations, the K readings made by the same observer are usually assumed to be true replications. In most situations, we use a general model $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$, $i = 1, \dots, n$, $j = 1, \dots, J$, $k = 1, \dots, K$ with the following minimal assumptions and notations: (1) μ_{ij} and ϵ_{ijk} are independent with means $E(\mu_{ij}) = \mu_j$ and $E(\epsilon_{ijk}) = 0$; (2) between-subject and within-subject variances $Var(\mu_{ij}) = \sigma_{Bj}^2$ and $Var(\epsilon_{ijk}) = \sigma_{Wj}^2$, respectively; (3) $Corr(\mu_{ij}, \mu_{ij'}) = \rho_{\mu ij'}$, $Corr(\mu_{ij}, \epsilon_{ij'k}) = 0$, $Corr(\epsilon_{ijk}, \epsilon_{ijk'}) = 0$ for all j, j', k, k' . Additional notations include $\sigma_j^2 = \sigma_{Bj}^2 + \sigma_{Wj}^2$, which is the total variability of observer j and $\rho_{jj'} = Corr(Y_{ijk}, Y_{ij'k})$ denotes the pairwise correlation between one reading from observer j and one reading from observer j' . In general, we have $\rho_{jj'} \leq \rho_{\mu ij'}$. If the K readings by an observer on a subject are not necessarily true replications, e.g., repeated measures over time, then additional structure may be needed for ϵ_{ijk} . In some specific situations, we may have $K = 1$ or $J = 2$, or ϵ_{ijk} may be decomposed further, with k being decomposed into multiple indices to denote multiple factors, such as time and treatment group.

3.1. Descriptive Tools

The basic descriptive statistics are the estimates of means μ_j , variances σ_j^2 , and correlation ρ_{jj} that can be obtained as sample mean, sample variance for each observer, and sample correlation between readings by any two observers. For data where the K readings by the same observer on a subject are true replications, one may obtain the estimates for between-subject variability σ_{Bj}^2 and within-subject variability σ_{Wj}^2 , $j = 1, \dots, J$ by fitting J one-way ANOVA models for each of the J observers. For data where k denotes the time of measurement, estimates for μ_j and σ_j^2 are obtained at each time point. These descriptive statistics provide intuition on how the J observers may deviate from each other on average, based on the first and second moments of the observers distribution. To help understand whether these values are statistically significant, confidence intervals should be provided along with the estimates. These descriptive statistics serve as an important intuitive component in understanding measurements made by observers. However, they do not fully quantify the degree of agreement between the J observers.

Several descriptive plots serve as visual tools in understanding and interpreting the data. These plots include (1) pairwise plots of any two readings, Y_{ijk} and $Y_{ij'k'}$ by observer j and j' on the n subjects with the 45 degree line as the reference line (Lin, 1989); (2) Bland and Altman plots (Bland and Altman, 1986) of average versus difference between any two readings by observer j and j' with the horizontal line of zero as the reference line. Both plots depict the visual examination of the overall agreement between observers j and j' . If k represents the time of reading, one may want to examine the plots at each time point.

3.2. Unscaled Agreement Indices

Summary agreement indices based on the absolute difference of readings by observers are grouped here as unscaled agreement indices. They are usually defined as the expectation of a function of the difference, or features of the distribution of the absolute difference. These indices include mean squared deviation, repeatability coefficient, repeatability variance, reproducibility variance (ISO), limits of agreement (Bland and Altman, 1999), coverage probability (CP), and total deviation index (TDI) (Choudhary, 2007a; Choudhary and Nagaraja, 2007; Lin et al., 2002).

3.2.1. Mean squared deviation, repeatability, and reproducibility. The *mean squared deviation (MSD)* is defined as the expectation of the squared difference of two readings. The *MSD* is usually used for the case of two observers, each making one reading for a subject ($K = 1$) (Lin et al., 2002). Thus

$$MSD_{jj'} = E(Y_{ij} - Y_{ij'})^2 = (\mu_j - \mu_{j'})^2 + (\sigma_j - \sigma_{j'})^2 + 2\sigma_j\sigma_{j'}(1 - \rho_{jj'}).$$

One should use an upper limit of *MSD* value, MSD_{ul} , to define satisfactory agreement as $MSD_{jj'} \leq MSD_{ul}$. In practice, MSD_{ul} may or may not be known; this can be a drawback to this measure of agreement. If d_0 is an acceptable difference between two readings, one may set d_0^2 as the upper limit.

Alternatively, it may be better to use $\sqrt{MSD_{jj'}}$ or $E(|Y_{ij} - Y_{ij'}|)$ than $MSD_{jj'}$ as a measure of agreement between observers j and j' , because one can interpret $\sqrt{MSD_{jj'}}$ or $E(|Y_{ij} - Y_{ij'}|)$ as the expected difference, and compare its value to d_0 , i.e., define $\sqrt{MSD_{jj'}} \leq d_0$ or $E(|Y_{ij} - Y_{ij'}|) \leq d_0$ as satisfactory agreement. It will be interesting to compare $\sqrt{MSD_{jj'}}$, $E(|Y_{ij} - Y_{ij'}|)$ to $MSD_{jj'}$ in simulation studies and in practical applications to examine their performance. Similarly, one may consider extending MSD by replacing the squared distance function with different distance functions (see examples in King and Chinchilli, 2001b for distance functions that are robust to the effects of outliers, or in Haber and Barnhart, 2007). It will be of interest to investigate the benefits of these possible new unscaled agreement indices.

The concept of MSD has been extended to the case of multiple observers, each taking multiple readings on a subject, where none of the observers is treated as a reference. Lin et al. (2007) defined overall, inter-, and intra- MSD for multiple observers under a two-way mixed model: $Y_{ijk} = \mu + \alpha_i + \beta_j + \gamma_{ij} + \epsilon_{ijk}$ where $\alpha_i \approx (0, \sigma_\alpha^2)$, (this notation means that α_i has mean 0 and variance of σ_α^2), $\gamma_{ij} \approx (0, \sigma_\gamma^2)$. Furthermore, the error term ϵ_{ijk} has mean 0 and a variance of σ_ϵ^2 . The observer effect β_j is assumed to be fixed with $\sum_j \beta_j = 0$, and we denote $\sigma_\beta^2 = \sum_j \sum_j (\beta_j - \beta_{j'})^2 / (J(J-1))$. The total, inter-, and intra- MSD are defined as: $MSD_{total}(Lin) = 2\sigma_\beta^2 + 2\sigma_\gamma^2 + 2\sigma_\epsilon^2$, $MSD_{inter}(Lin) = 2\sigma_\beta^2 + 2\sigma_\gamma^2 + 2\sigma_\epsilon^2/K$, $MSD_{intra}(Lin) = 2\sigma_\epsilon^2$. The above definitions require equal variance assumptions in the two-way mixed model. One extension is to define the MSD for multiple observers without any assumptions such as:

$$MSD_{total} = \frac{\sum_{j=1}^J \sum_{j'=j+1}^J \sum_{k=1}^K \sum_{k'=1}^K E(Y_{ijk} - Y_{ijk'})^2}{J(J-1)K^2}$$

$$MSD_{inter} = \frac{\sum_{j=1}^J \sum_{j'=j+1}^J E(\mu_{ij} - \mu_{ij'})^2}{J(J-1)}$$

$$MSD_{intra} = \frac{\sum_{j=1}^J \sum_{k=1}^K \sum_{k'=k+1}^K E(Y_{ijk} - Y_{ijk'})^2}{JK(K-1)},$$

where $\mu_{ij} = E(Y_{ijk})$. One can also define $MSD_{j,intra}$ for each observer as

$$MSD_{j,intra} = \frac{\sum_{k=1}^K \sum_{k'=k+1}^K E(Y_{ijk} - Y_{ijk'})^2}{K(K-1)}.$$

Thus, MSD_{intra} is the average of J separate $MSD_{j,intra}$'s. Although the above definition involves k , the $MSDs$ do not depend on K as long as we have true replications. For these general definitions of the $MSDs$, one can show that $MSD_{total} = MSD_{inter} + MSD_{intra}$ (Haber et al., 2005). We note that under the two-way mixed model, the general MSD_{total} and MSD_{intra} reduce to the $MSD_{total}(Lin)$ and $MSD_{intra}(Lin)$, respectively and the general MSD_{inter} is the limit of Lin's $MSD_{inter}(Lin)$ as $K \rightarrow \infty$. It would be of interest to write down the expressions of these $MSDs$ based on the general model, $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$ in place of the two-way mixed model. One can

also extend the total MSD and inter- MSD to the case where the J th observer is treated as a reference by using

$$MSD_{total}^R = \frac{\sum_{j=1}^{J-1} \sum_{k=1}^K \sum_{k'=1}^K E(Y_{ijk} - Y_{ijk'})^2}{(J-1)K^2}$$

$$MSD_{inter}^R = \frac{\sum_{j=1}^{J-1} E(\mu_{ij} - \mu_{iJ})^2}{J-1}.$$

Pairwise $MSDs$ may be used to model these $MSDs$ as a function of the observers' characteristics, e.g., experienced or not. If a subject's covariates are available, one may be also interested in modeling the MSD as a function of covariates such as age, gender, etc., by specifying the logarithm of the MSD (because the MSD is positive in general) as a linear function of covariates.

Repeatability standard deviation (ISO, 1994) is defined as the dispersion of the distribution of measurements under repeatability conditions. Thus, *repeatability standard deviation* for observer j is defined as σ_{w_j} and *repeatability variance* for observer j is $\sigma_{w_j}^2$. *Repeatability coefficient* or *repeatability limit* for observer j is $1.96\sqrt{2\sigma_{w_j}^2}$, which is interpreted as the value within which any two readings by the j th observer would lie for 95% of subjects (Bland and Altman, 1999; ISO, 1994). It is expected that the repeatability variances are different for different observers. The ISO assumed that such differences are small, and stated that it is justifiable to use a common value for the overall repeatability variance as

$$\sigma_r^2 = \frac{\sum_{j=1}^J \sigma_{w_j}^2}{J}.$$

In this case, we have $\sigma_r^2 = MSD_{intra}/2$. Therefore, one can define the overall repeatability standard deviation, repeatability coefficient, or limit for J observers by using this σ_r^2 with the additional assumptions.

Reproducibility standard deviation (ISO, 1994) is defined as the dispersion of the distribution of measurements under reproducibility conditions. If the reproducibility condition is the usage of J observers, the ISO 5725-1 used the one-way model $Y_{ijk} = \mu + \beta_j + \epsilon_{ijk}$ to define *reproducibility variance* as

$$\sigma_R^2 = \sigma_\beta^2 + \sigma_r^2.$$

where $\sigma_\beta^2 = Var(\beta_j)$ with observers treated as random. One may use the notation $\sigma_\beta^2 = \sum_{j=1}^J (\beta_j - \beta_\bullet)^2 / (J-1)$ for the above formula if observers are treated as fixed with $\beta_\bullet = \sum_{j=1}^J \beta_j / J$. In this case, we note that $\sigma_R^2 = MSD_{total}/2$. The reproducibility standard deviation is the square root of σ_R^2 and the *reproducibility coefficient* or *reproducibility limit* is $1.96\sqrt{2\sigma_R^2}$.

We see that the repeatability variance and reproducibility variance correspond to half of the MSD_{intra} and half of MSD_{total} , respectively, under the assumptions used in the ISO. This suggests that one can extend the definition of repeatability and reproducibility variances more generally as

$$\sigma_r^2 = MSD_{intra}/2, \quad \sigma_R^2 = MSD_{total}/2$$

by utilizing the general definitions of the *MSDs* without making the assumptions used in the ISO.

Estimations for *MSD*, repeatability, and reproducibility are usually carried out by using the sample counterparts of the means and variance components. Inference is often based on large sample distribution of these estimates or based on normal assumptions regarding distribution of the data.

It is clear that *MSD*, reproducibility standard deviation, reproducibility variance, and reproducibility coefficient are measures of agreement between observers. They are also measures of validity between two observers if one of them is the reference standard. Repeatability standard deviation, repeatability variance, and repeatability coefficient are measures of agreement between replicated readings within observers, and thus, measures of precision.

3.2.2. Limits of agreement. Due to simplicity and intuitive appeal, limits of agreement (LOA) (Altman and Bland, 1983, 1986, 1999; Bland and Altman, 2007) are widely used (Bland and Altman, 1992; Ryan and Woodall, 2005) for assessing agreement between two observers in medical literature. This method was developed for two observers that can then be used for pairwise comparisons of J observers. The key principle of the LOA method is to estimate the difference of single observations between two observers and the corresponding $(1 - \alpha)$ 100% probability interval (PI) that contains middle $1 - \alpha$ probability of the distribution of difference. The estimates for the limits of this PI are called the limits of agreement. Let $D_i = Y_{i1} - Y_{i2}$ be the difference between the single observations of the two observers. The 95% LOA are $\mu_D \pm 1.96\sigma_D$ where $\mu_D = E(D_i)$ and $\sigma_D^2 = \text{Var}(D_i)$, under normality assumption on D_i . If the absolute limit is less than an acceptable difference, d_0 , then the agreement between the two observers is deemed satisfactory.

For data without replications, the LOA can be estimated by replacing μ_D and σ_D^2 by the sample mean, D_\bullet , and sample variance, S_D^2 , of D_i . The variances of these estimated limits are $(1/n \pm 1.96^2/(2(n-1)))\sigma_D^2$ that can be estimated by replacing σ_D^2 by the sample variance of D_i (Bland and Altman, 1999). The key implicit assumption for the method of estimation is that the difference between the two observers is reasonably stable across the range of measurements. Bland and Altman also showed how to compute the LOA if μ_D depends on a covariate or σ_D^2 depends on the average readings. The LOA is often displayed in the popular Bland and Altman plot (average, $(Y_{i1} + Y_{i2})/2$, versus difference, $Y_{i1} - Y_{i2}$) with two horizontal lines of the estimated LOA: $D_\bullet \pm 1.96S_D$ and two horizontal lines of the 95% lower bound of the lower limit and 95% upper bound of the upper limit:

$$D_\bullet - 1.96S_D - 1.96\sqrt{\left(\frac{1}{n} - \frac{1.96^2}{2(n-1)}\right)S_D^2}, \quad D_\bullet + 1.96S_D + 1.96\sqrt{\left(\frac{1}{n} + \frac{1.96^2}{2(n-1)}\right)S_D^2}.$$

Lin et al. (1998) argued that instead of using the above two-sided CIs for the estimated LOA, one can use a one-sided upper confidence bound (UCB) for $\mu_D + 1.96\sigma_D$ and a one-sided lower confidence bound (LCB) for $\mu_D - 1.96\sigma_D$ to derive $UCB = D_\bullet + a_n S_D$ and $LCB = D_\bullet - a_n S_D$ with $a_n = 1.96 + 1.71n^{-1/2}t_{n-1}(\alpha)$ where $t_{n-1}(\alpha)$ is the upper α th percentile of a t_k distribution. Interval (LCB, UCB) is closely related to the tolerance interval (Choudhary, 2007a,b; Choudhary and Nagaraja, 2007) for $TDI_{0.95}$, an index discussed in Section 3.2.3.

The LOA method has also been extended to data with replications or with repeated measures (Bland and Altman, 1999, 2007). Bland and Altman (2007) distinguish two situations: (1) multiple time-matched observations per individual by two observers where the true value of the subject may or may not change over time; (2) multiple observations per individual (not time-matched) by two observers where the true value of the subject is constant at a prespecified length of time when the two observers take measurements. Naturally the LOA in the first situation is narrower than in the second because of reduced variability due to time-matching. One can think of the first situation as time-matched repeated measures of observers on the same subject and the second situation as unmatched replications by observers on the same subject. The LOAs for these two situations are still defined as $\mu_D \pm 1.96\sigma_D$, except that now D_i is defined differently and thus the LOAs for these two situations have slightly different interpretations. In the first situation, $D_{ik} = Y_{i1k} - Y_{i2k}$, $i = 1, \dots, n$, $k = 1, \dots, K_i$ with $\mu_D = E(D_{ik})$ and $\sigma_D^2 = \text{Var}(D_{ik})$ for all k , where Y_{i1k} and Y_{i2k} are two observations made by the two observers at the same time. The underlying true value of the subject may or may not change over time when K_i measurements are taken by an observer. This may correspond to the cases of matched repeated measures or matched replications, respectively. Note that if $K_i = 1$ for all i , it reduces to the original situation of no multiple observations.

In the second situation, $D_{ik_i k'_i} = Y_{i1k_i} - Y_{i2k'_i}$, $i = 1, \dots, n$, $k_i = 1, \dots, K_{1i}$, $k'_i = 1, \dots, K_{2i}$ with $\mu_D = E(D_{ik_i k'_i})$ and $\sigma_D^2 = \text{Var}(D_{ik_i k'_i})$ for all k_i and k'_i where Y_{i1k_i} and $Y_{i2k'_i}$ are two observations made by the two observers at any time of a specified interval (e.g., 1 day) within which the underlying true value of the subject does not change. The number of observations on a subject may differ for different observers. Due to time matching in the first situation, one would expect that the σ_D^2 in the first situation is smaller than the one from the second situation. However, if there is no time effect in the second situation, one would expect these two variances to be similar. The LOA in the first situation has the interpretation of the LOA between two observers who made single observations at the same time. The LOA in the second situation has the interpretation of limits of agreement between two observers who made single observations at a specified time interval. We emphasize that the focus is still on the LOA for single observations between two observers, although multiple observations per observer are used to obtain estimates for μ_D and σ_D^2 .

Bland and Altman (1999, 2007) described a method of moment approach to estimating μ_D and σ_D^2 for data with multiple observations in both situations. In both situations, μ_D is estimated as $\hat{\mu}_D = Y_{\bullet 1\bullet} - Y_{\bullet 2\bullet}$ averaging over indices of i and k or k' . In the first situation, σ_D^2 is estimated via a one-way ANOVA model for D_{ik} ,

$$D_{ik} = \mu_D + I_{Di} + E_{Dik},$$

where I_{Di} represents the subject by observer interaction, and E_{Dij} represents independent random error within the subject for that pair of observations. The implicit assumption for this model is that there is no time effect in D_{ik} , even though there may be a time effect in Y_{ijk} . In other words, D_{ik} 's are treated as replications of $\mu_D + I_{Di}$. Under this model, $\sigma_D^2 = \sigma_{DI}^2 + \sigma_{DW}^2$ where $\sigma_{DI}^2 = \text{Var}(I_{Di})$ and $\sigma_{DW}^2 = \text{Var}(E_{Dik})$. Let MSB_D and MSW_D be the between-subject and within-subject

mean sums of squares from this one-way ANOVA model. Then the method of moment estimator for σ_D^2 is

$$\hat{\sigma}_D^2 = \frac{MSB_D - MSW_D}{\frac{(\sum K_i)^2 - \sum K_i^2}{(n-1) \sum K_i}} + MSW_D,$$

where the first term on the right estimates σ_{DI}^2 and the second term estimates σ_{DW}^2 . In the second situation, two one-way ANOVA models are used, one for each observer:

$$Y_{i1k_i} = \mu_1 + I_{i1} + E_{i1k_i}, \quad k_i = 1, \dots, K_{1i}, \quad Y_{i2k'_i} = \mu_1 + I_{i1} + E_{i2k'_i}, \quad k'_i = 1, \dots, K_{2i}.$$

Thus, $\sigma_D^2 = \text{Var}(Y_{i1k_i} Y_{i2k'_i}) = \sigma_{I1}^2 + \sigma_{I2}^2 + \sigma_{W1}^2 + \sigma_{W2}^2$, where $\sigma_{Ij}^2 = \text{Var}(I_{ij})$, $\sigma_{Wj}^2 = \text{Var}(E_{ijk_i})$, $j = 1, 2$. The implicit assumption is that Y_{ijk_i} 's, $k_i = 1, \dots, K_i$ are replications of $\mu_j + I_{ij}$ given i and j . Note that $\text{Var}(Y_{\bullet 1\bullet} - Y_{\bullet 2\bullet}) = \sigma_{I1}^2 + \frac{1}{n}(\sum \frac{1}{K_{1i}})\sigma_{W1}^2 + \sigma_{I2}^2 + \frac{1}{n}(\sum \frac{1}{K_{2i}})\sigma_{W2}^2$.

Thus,

$$\sigma_D^2 = \text{Var}(Y_{\bullet 1\bullet} - Y_{\bullet 2\bullet}) + \left(1 - \frac{1}{n} \left(\sum \frac{1}{K_{1i}}\right)\right) \sigma_{W1}^2 + \left(1 - \frac{1}{n} \left(\sum \frac{1}{K_{2i}}\right)\right) \sigma_{W2}^2.$$

which can be estimated by replacing $\text{Var}(Y_{\bullet 1\bullet} - Y_{\bullet 2\bullet})$ by its sample variance from data $(Y_{i1\bullet} - Y_{i2\bullet})$, $i = 1, \dots, n$ and replacing σ_{Wj}^2 by MSW_j , the within-subject mean sums of squares from the one-way ANOVA model for observer j . It will be of interest to extend the LOA for more than two observers.

3.2.3. Coverage probability and total deviation index. As elaborated by Lin and colleagues (Lin, 2000a,b; Lin et al., 2002), an intuitive measure of agreement is a measure that captures a large proportion of data within a boundary for allowed observers' differences. The proportion and boundary are two quantities that correspond to each other. If we set d_0 as the predetermined boundary; i.e., the maximum acceptable absolute difference between two observers' readings, we can compute the probability of absolute difference between any two observers' readings less than d_0 . This probability is called *coverage probability (CP)*. On the other hand, if we set π_0 as the predetermined coverage probability, we can find the boundary so that the probability of absolute difference less than this boundary is π_0 . This boundary is called *total deviation index (TDI)* and is the $100\pi_0$ percentile of the absolute difference of paired observations. A satisfactory agreement may require a large *CP* or, equivalently, a small *TDI*. For $J = 2$ observers, let Y_{i1} and Y_{i2} be the readings of these two observers, the *CP* and *TDI* are defined as

$$CP_{d_0} = \text{Prob}(|Y_{i1} - Y_{i2}| < d_0), \quad TDI_{\pi_0} = f^{-1}(\pi_0)$$

where $f^{-1}(\pi_0)$ is the solution of d by setting $f(d) = \text{Prob}(|Y_{i1} - Y_{i2}| < d) = \pi_0$.

Estimation and inference on CP_{d_0} and TDI_{π_0} often requires a normality assumption on $D_i = Y_{i1} - Y_{i2}$. Assume that D_i is normally distributed with mean μ_D

and variance σ_D^2 . We have

$$CP_{d_0} = \Phi\left(\frac{d_0 - \mu_D}{\sigma_D}\right) - \Phi\left(\frac{-d_0 - \mu_D}{\sigma_D}\right) = \chi_1^2\left(d_0^2, \frac{\mu_D^2}{\sigma_D^2}\right) \approx CP_{d_0}^* = \chi_1^2\left(\frac{d_0^2}{MSD_{12}}\right)$$

$$TDI_{\pi_0} = \sigma_d \sqrt{\chi_1^{2(-1)}\left(\pi_0, \frac{\mu_D^2}{\sigma_D^2}\right)} \approx TDI_{\pi_0}^* = Q_0(\mu_D^2 + \sigma_D^2) = Q_0 \sqrt{MSD_{12}},$$

where $\Phi(t)$ is the cumulative distribution function of standard normal distribution, $MSD_{12} = E(Y_{i1} - Y_{i2})^2 = \mu_D^2 + \sigma_D^2$ is the MSD between the two observers, $\chi_1^2(t)$ is the cumulative distribution of chi-square distribution with one degree of freedom, $\chi_1^{2(-1)}(\pi_0, \lambda)$ is the inverse of the chi-square distribution with one degree of freedom and the non-centrality parameter λ , and $Q_0 = \Phi^{-1}\left(\frac{1+\pi_0}{2}\right)$ with $\Phi^{-1}(t)$ as the inverse function of $\Phi(t)$. Point estimation for CP_{d_0} is obtained by plugging the sample counterparts for μ_D and σ_D^2 ; the inference is based on the asymptotic distribution for $\ln(\widehat{CP}_{d_0}/(1 - \widehat{CP}_{d_0}))$ described in Lin et al. (2002).

Similarly, estimation for TDI_{π_0} is obtained by plugging the sample counterparts of the parameters; the inference is based on approximation for TDI_{π_0} with $2 \ln(TDI_{\pi_0}^*) = 2 \ln Q_0 + \ln MSD_{12}$ by using the asymptotic distribution of $\ln(\widehat{MSD}_{12})$ (Lin et al., 2002). This approximation is usually reasonable if μ_D^2/σ_D^2 is small. Due to approximation, different conclusions may be reached with the above method of inference, especially for small sample sizes and large values of μ_D^2/σ_D^2 when testing the following two equivalent hypotheses:

$$H_0 : CP_{d_0} \leq \pi_0 \quad vs \quad H_1 : CP_{d_0} > \pi_0; \quad (1)$$

$$H_0 : TDI_{\pi_0} \geq d_0 \quad vs \quad H_1 : TDI_{\pi_0} < d_0. \quad (2)$$

Alternative inference approaches are available in this situation. Wang and Hwang (2001) proposed a nearly unbiased test (NUT) for testing (1) and Choudhary and Nagaraja (2007) proposed an exact test and modified NUT for testing (1) and equivalently for testing (2) for data with a small sample size (≤ 30) and a bootstrap test for data with a moderate sample size. These tests appear to outperform previous tests in terms of maintaining the type I error rate close to the nominal level for all combinations of parameter values under the normality assumption. Rather than computing a $(1 - \alpha)100\%$ confidence interval on an approximated value of $TDI_{\pi_0}^*$, they provided a $1 - \alpha$ upper confidence bound, U , for TDI_{π_0} and used $(-U, U)$ as a $1 - \alpha$ confidence tolerance interval with a probability content of π_0 for the distribution of difference. An interval (L, U) is a tolerance interval with a probability content p and confidence $1 - \alpha$ if $P(F(U) - F(L) \geq p) = 1 - \alpha$, where F is the cumulative distribution function of X (Guttman, 1988). For $\pi_0 = 0.95$ and $\alpha = 0.05$, this tolerance interval is expected to be wider than one based on LOAs, because the probability content of the tolerance interval is 0.95 with confidence level 95%, whereas the probability content of the 95% LOAs is approximately 0.95 on average. Choudhary and Nagaraja (2007) extended their approach to incorporate a continuous covariate (Choudhary, 2007c; Choudhary and Ng, 2006) and to deal with data with replications and longitudinal measurements (Choudhary, 2007a).

Lin et al. (2007) extended the concept of *CP* and *TDI* to data with more than two observers making multiple observations on a subject, where none of the observers is treated as reference. Using the approximations that link the *CP* and the *TDI* with the *MSD*, they define *CP* and *TDI* for multiple observers as

$$CP_{d_0total} = \chi_1^2 \left(\frac{d_0^2}{MSD_{total}} \right), \quad CP_{d_0inter} = \chi_1^2 \left(\frac{d_0^2}{MSD_{inter}} \right), \quad CP_{d_0intra} = \chi_1^2 \left(\frac{d_0^2}{MSD_{intra}} \right)$$

$$TDI_{\pi_0total} = Q_0 \sqrt{MSD_{total}}, \quad TDI_{\pi_0inter} = Q_0 \sqrt{MSD_{inter}}, \quad TDI_{\pi_0intra} = Q_0 \sqrt{MSD_{intra}}$$

where MSD_{total} , MSD_{inter} , MSD_{intra} were defined in Section 3.2.1 under the two-way mixed model with normality assumption. One can extend the *CP* and *TDI* to the case where the J th observer is treated as a reference by using the *MSD* for the case of the J th observer as a reference.

3.2.4. Comments. In summary, the unscaled agreement indices of *MSD*, *LOA*, *CP*, and *TDI* are all based on the differences between observers' readings for the case that none of the observers is treated as a reference. These indices are related to each other under some assumptions. First, *MSD* has approximately one-to-one correspondence to the *CP* and *TDI* under normality assumption and small value for μ_D^2/σ_D^2 . Therefore, one has the same criterion of agreement based on *MSD*, *CP*, or *TDI* indices. If we use an upper limit, MSD_{ul} for declaring satisfactory agreement, i.e., $MSD < MSD_{ul}$, this should correspond to using $P_{d_0} > \chi_1^2 \left(\frac{d_0^2}{MSD_{ul}} \right)$ or $TDI_{\pi_0} < Q_0 \sqrt{MSD_{ul}}$ for declaring satisfactory agreement.

Second, for two observers who do not make replicated measurements, the *MSD* and *LOA* are related by $LOA = \mu_D \pm 1.96 \sqrt{MSD_{12} - \mu_D^2}$, because $MSD_{12} = \mu_D^2 + \sigma_D^2$. In particular, if there are no systematic mean shifts (i.e., $\mu_j = \mu$ for $j = 1, 2$) then the 95% *LOA* corresponds to $\pm 1.96 \sqrt{MSD_{total}}$, whose absolute value is the 95% reproducibility limit. For data with replicated measurements with two observers, this relationship holds by replacing MSD_{12} by MSD_{total} . Third, a tolerance interval derived using $TDI_{0.95}$ and the 95% *LOA* have related interpretations. It will be of interest to compare these two intervals under normality assumption. Future research is needed for these unscaled agreement indices when one of the observers is treated as a reference. Also of interest is the behavior of these indices for non-normal data.

3.3. Scaled Agreement Indices

3.3.1. Intraclass correlation coefficient. Historically, agreement between quantitative measurements has been evaluated via the intraclass correlation coefficient (ICC). Numerous versions of ICC (Bartko, 1966, 1974; Eliasziw et al., 1994; Müller and Büttner, 1994; McGraw and Wong, 1996; Shrout and Fleiss, 1979) have been proposed in many areas of research by assuming different underlying ANOVA models for the situation where none of the observers is treated as reference. In earlier research on ICCs (prior to McGraw and Wong's work in 1996), ICCs were rigorously defined as the correlation between observations from different observers under different ANOVA model assumptions where observers are treated as random. An ICC (denoted as ICC_{3c} below) is also proposed under the two-way mixed model where the observers are treated as fixed, using the concept of

correlation. The ICC_{3c} was originally proposed by Bartko (1966) and later corrected by Bartko (1974) and advocated by Shrout and Fleiss (1979). McGraw and Wong (1996) suggested calling ICC_{3c} as ICC for consistency and proposed an ICC for agreement (denoted here as ICC_3). They also added an ICC for the two-way mixed model without interaction (denoted here as ICC_2).

We unite different versions of ICCs into *three* ICCs under three kinds of model assumption with unifying notations for both cases of random and fixed observers. We do not present ICCs for averaged observations (McGraw and Wong, 1996; Shrout and Fleiss, 1979), as we are interested in agreement between single observations. We assume that each observer takes K readings on each subject where $K = 1$ if there is no replication and $K \geq 2$ if there are replications (Eliaszew et al., 1994). We discuss situations of $K = 1$ and $K \geq 2$ when comparing ICC to CCC in Section 3.3.1. For each ICC, estimates are obtained via method of moment based on the expectation of the mean sums of squares from these different ANOVA models. The definitions of the three types of ICCs and their corresponding estimates are presented below:

- ICC_1 is based on a one-way random effect model without observer effect (Bartko, 1966; McGraw and Wong, 1996; Shrout and Fleiss, 1979):

$$Y_{ijk} = \mu + \alpha_i + \epsilon_{ijk}$$

with assumptions: $\alpha_i \sim N(0, \sigma_\alpha^2)$; $\epsilon_{ijk} \sim N(0, \sigma_\epsilon^2)$; and ϵ_{ijk} is independent of α_i .

$$ICC_1 = \frac{\sigma_\alpha^2}{\sigma_\alpha^2 + \sigma_\epsilon^2}, \quad \widehat{ICC}_1 = \frac{MS_\alpha - MS_\epsilon}{MS_\alpha + (JK - 1)MS_\epsilon},$$

where MS_α and MS_ϵ are the mean sums of squares from the one-way ANOVA model for between and within subjects, respectively.

- ICC_2 is based on a two-way mixed or random (depending on whether the observers are fixed or random) effect model without the observer-subject interaction (McGraw and Wong, 1996):

$$Y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon_{ijk}$$

with assumptions: $\alpha_i \sim N(0, \sigma_\alpha^2)$, $\epsilon_{ijk} \sim N(0, \sigma_\epsilon^2)$, and ϵ_{ijk} is independent of α_i . β_j is treated as either as a fixed or a random effect, depending on whether the observers are fixed or random. If observers are fixed, notation $\sigma_\beta^2 = \sum_{j=1}^J \beta_j^2 / (j - 1)$ is used with constraint of $\sum_{j=1}^J \beta_j = 0$. If observers are random, additional assumptions are $\beta_j \sim N(0, \sigma_\beta^2)$ and $\alpha_i, \beta_j, \epsilon_{ijk}$ are mutually independent.

$$ICC_2 = \frac{\sigma_\alpha^2}{\sigma_\alpha^2 + \sigma_\beta^2 + \sigma_\epsilon^2}, \quad \widehat{ICC}_2 = \frac{MS_\alpha - MS_\epsilon}{MS_\alpha + (JK - 1)MS_\epsilon + J(MS_\beta - MS_\epsilon)/n}.$$

- ICC_3 is based on a two-way mixed or random effect model (depending on whether the observers are fixed or random) with observer-subject interaction (Eliaszew et al., 1994; McGraw and Wong, 1996).

$$Y_{ijk} = \mu + \alpha_i + \beta_j + \gamma_{ij} + \epsilon_{ijk}$$

with assumptions: $\alpha_i \sim N(0, \sigma_\alpha^2)$, $\epsilon_{ijk} \sim N(0, \sigma_\epsilon^2)$, and ϵ_{ijk} is independent of α_i . If observers are fixed, notation $\sigma_\beta^2 = \sum_{j=1}^J \beta_j^2 / (J-1)$ is used with constraint of $\sum_{j=1}^J \beta_j = 0$ and $\gamma_{ij} \sim N(0, \sigma_\gamma^2)$. If the observers are random, additional assumptions are $\beta_j \sim N(0, \sigma_\beta^2)$, $\gamma_{ij} \sim N(0, \sigma_\gamma^2)$ and $\alpha_i, \beta_j, \gamma_{ij}, \epsilon_{ijk}$ are mutually independent.

$$ICC_3 \text{ (fixed } \beta_j) = \frac{\sigma_\alpha^2 - \sigma_\gamma^2 / (J-1)}{\sigma_\alpha^2 + \sigma_\beta^2 + \sigma_\gamma^2 + \sigma_\epsilon^2}, \quad ICC_3 \text{ (random } \beta_j) = \frac{\sigma_\alpha^2}{\sigma_\alpha^2 + \sigma_\beta^2 + \sigma_\gamma^2 + \sigma_\epsilon^2}$$

$$\widehat{ICC}_3 = \frac{MS_\alpha - MS_\gamma}{MS_\alpha + J(K-1)MS_\epsilon + (J-1)MS_\gamma + J(MS_\beta - MS_\gamma)/n}.$$

As mentioned earlier, Bartko (1974) and Shrout and Fleiss (1979) presented the following ICC, later called ICC for consistency (e.g., observers are allowed to differ by a fixed constant) by McGraw and Wong (1996) using

$$ICC_{3c} = \frac{\sigma_\alpha^2 - \sigma_\gamma^2 / (J-1)}{\sigma_\alpha^2 + \sigma_\gamma^2 + \sigma_\epsilon^2}$$

when observers are treated as a fixed effect. The ICC_{3c} does not take into account systematic shifts by observers in the denominator; thus, it is measure of consistency rather than agreement. For all three ICCs, if the observer is a random effect, the ICC defined with the assumed ANOVA model is equal to $corr(Y_{ijk}, Y_{ij'k'})$. If the observer is a fixed effect, we have $corr(Y_{ijk}, Y_{ij'k'}) = ICC_{3c}$ under the two-way mixed model with interaction. We also note that if there is no replication ($K=1$), we have $MS_\gamma = MS_\epsilon$, which estimates $\sigma_\gamma^2 + \sigma_\epsilon^2$ in the expression for \widehat{ICC}_3 . Thus, for $K=1$, we have $\widehat{ICC}_2 = \widehat{ICC}_3$. Inference about ICCs is well developed and McGraw and Wong (1996) provided a very detailed summary. All ICCs require an assumption of normality, equal variances of $Var(Y_{ijk} | j)$, and equal pairwise correlations of $Corr(Y_{ijk}, Y_{ij'k'})$. The ICC_1 also requires the additional assumption of equal means of $E(Y_{ijk} | j)$ and corresponds to the original ICC that requires measurements to be parallel (see Section 2.3). The assumptions for ICC_1 may be reasonable if there is only one observer taking replicated measurements. In this case, ICC_1 assesses test-retest reliability, or scaled repeatability. With additional assumptions, ICC_3 reduces to ICC_2 and ICC_2 reduces to ICC_1 .

The assumptions used to define ICC are the main disadvantages to using ICCs to assess agreement. Of note is the fact that all ICCs are increasing functions of between-subject variability (represented here by σ_α^2). Thus, it would attain a high value for a population with substantial heterogeneity. Due to this fact, Vangeneugden et al. (2004, 2005) and Molenberghs et al. (2007) interpret the ICC as a reliability measure that assesses the degree of differentiation of subjects from a population, rather than agreement.

The ICCs presented here may be used for data with repeated measures where k denotes the time of the measurement. However, these ICCs may not be very useful unless one modifies the assumptions on ϵ_{ijk} in order to take into account the time structure. A linear mixed-model approach to estimate reliability

for repeated measures has been proposed by Vangeneugden et al. (2004) and Molenberghs et al. (2007).

3.3.2. Concordance correlation coefficient. The CCC is the most popular index for assessing agreement in the statistical literature. The CCC was originally developed by Lin (1989) for two observers ($J = 2$), each making a single reading on a subject. It was later extended to multiple J observers for data without replications (Barnhart et al., 2002; King and Chinchilli, 2001a; Lin, 1989; Lin et al., 2002) and for data with replications (Barnhart et al., 2005a; Lin et al., 2007) where none of the observers is treated as reference. These extensions included the original CCC for $J = 2$ as a special case. Recently, Barnhart et al. (2007b) extended the CCC to the situation where one of the multiple observers is treated as reference. Barnhart and Williamson (2001) used a generalized estimating equations (GEE) approach to modeling pairwise CCCs as a function of covariates. Chinchilli et al. (1996) and King et al. (2007a,b) extended the CCC for data with repeated measures comparing two observers. Quiroz (2005) extends the CCC for data with repeated measures comparing multiple observers by using the two-way ANOVA model without interaction. Due to the assumptions of the ANOVA model, the CCC defined by Quiroz (2005) is the special case of CCC by Barnhart et al. (2005a) for data with replications.

We first present the total CCC, inter-CCC, and intra-CCC for multiple observers for data with replications where none of the observers is treated as a reference. Of note here is the fact that the total CCC is the usual CCC for data without replications. The definition of the total CCC does not require replicated data (Barnhart et al., 2002; King and Chinchilli, 2001a), although one can estimate both the between-subject (σ_{Bj}^2) and within-subject (σ_{Wj}^2) variabilities for data with replications, while only total variability ($\sigma_j^2 = \sigma_{Bj}^2 + \sigma_{Wj}^2$) can be estimated for data without replications. One cannot estimate inter- or intra-CCCs for data without replications.

Let Y_{ijk} be the k th replicated measurements for the i th subject by the j th method and write $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$ with the assumptions and notations in the beginning of Section 3. The CCC for assessing agreement between J observers using data with or without replications can be written

$$CCC_{total} = \rho_c = 1 - \frac{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J E(Y_{ijk} - Y_{ij'k'})^2}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J E(Y_{ijk} - Y_{ij'k'})^2} \quad (3)$$

$$= \frac{2 \sum_{j=1}^{J-1} \sum_{j'=j+1}^J \sigma_j \sigma_{j'} \rho_{jj'}}{(J-1) \sum_{j=1}^J \sigma_j^2 + \sum_{j=1}^{J-1} \sum_{j'=j+1}^J (\mu_j - \mu_{j'})^2} \quad (4)$$

$$= \frac{2 \sum_{j=1}^{J-1} \sum_{j'=j+1}^J \sigma_{Bj} \sigma_{Bj'} \rho_{jj'}}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J [2\sigma_{Bj} \sigma_{Bj'} + (\mu_j - \mu_{j'})^2 + (\sigma_{Bj} - \sigma_{Bj'})^2 + \sigma_{Wj}^2 + \sigma_{Wj'}^2]} \quad (5)$$

where E_j is the conditional expectation given independence of Y_{ijk} , $Y_{ij'k'}$. Although the above definition involves k , the CCC does not depend on K as long as they are true replications. Expression (4) can be used for the CCC for data without

replications, while both expressions (4) and (5) can be used for total CCC for data with replications. With replications, this CCC is the total CCC defined in Barnhart et al. (2005a) and Lin et al. (2007). Fay (2005) called the CCC a fixed marginal agreement coefficient (FMAC) and proposed a random marginal agreement coefficient (RMAC) by replacing $E_I(Y_{ijk} - Y_{ij'k'})^2$ with $E_{Z_j} E_{Z_{j'}}(Z_j - Z_{j'})^2$, where Z_j and $Z_{j'}$ are independent and identically-distributed random variables with a mixture distribution of random variable, $0.5Y_j + 0.5Y_{j'}$. We note later in a comparison of CCC and ICC that this RMAC is closely related to $E(\widehat{ICC}_1)$ with expectations taken under the general model of $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$. Barnhart et al. (2005a) also defined inter-CCC at the level of μ_{ij} 's as

$$\begin{aligned} CCC_{inter} = \rho_c(\mu) &= 1 - \frac{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J E(\mu_{ij} - \mu_{ij'})^2}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J E_I(\mu_{ij} - \mu_{ij'})^2} \\ &= \frac{2 \sum_{j=1}^{J-1} \sum_{j'=j+1}^J \sigma_{Bj} \sigma_{Bj'} \rho_{\mu jj'}}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J [2\sigma_{Bj} \sigma_{Bj'} + (\mu_j - \mu_{j'})^2 + (\sigma_{Bj} - \sigma_{Bj'})^2]} \end{aligned}$$

and intra-CCC for observer j as

$$\begin{aligned} CCC_{j,intra} = \rho_j^I &= 1 - \frac{\sum_{k=1}^{K-1} \sum_{k'=k+1}^J E(Y_{ijk} - Y_{ijk'})^2}{\sum_{k=1}^{K-1} \sum_{k'=k+1}^J E_I(Y_{ijk} - Y_{ijk'})^2} \\ &= ICC_{1j} = \frac{\sigma_{Bj}^2}{\sigma_{Bj}^2 + \sigma_{Wj}^2}, \end{aligned}$$

which is the ICC_1 for observer j in Section 3.3.1. The total CCC, inter-CCC, and intra-CCCs are related by

$$\frac{1}{\rho_c} = \frac{1}{\rho_c(\mu)} + \frac{1}{\gamma},$$

where

$$\frac{1}{\gamma} = \frac{(J-1) \sum_{j=1}^J \sigma_{Wj}^2}{2 \sum_{j=1}^{J-1} \sum_{j'=j+1}^J \sigma_{Bj} \sigma_{Bj'} \rho_{\mu jj'}} = \sum_{j=1}^J \omega_j \frac{1 - \rho_j^I}{\rho_j^I}$$

with $\omega_j = \sigma_{Bj}^2 / (2 \sum_{j=1}^{J-1} \sum_{j'=j+1}^J \sigma_{Bj} \sigma_{Bj'} \rho_{\mu jj'})$ is the weighted sum of the odds of $1 - \rho_j^I$.

Lin et al. (2007) defined an inter-CCC, $CCC_{inter}(\text{Lin})$, at the level of average readings $Y_{ij\bullet}$, rather than at the level of μ_{ij} . Thus, Lin's inter-CCC depends on the number of replications; as the number of replications approaches infinity, it becomes the inter-CCC defined by Barnhart et al. (2005a), (i.e., $CCC_{inter}(\text{Lin}) \rightarrow CCC_{inter}$ as $K \rightarrow \infty$). Lin et al. (2007) also define an overall intra-CCC, rather than separate intra-CCCs for each observer. This overall intra-CCC is the average of the J intra-CCCs above.

Barnhart et al. (2007a,b) extended these CCCs to the case where the J th observer is treated as a reference. In this case, while definition for the intra-CCCs

remains the same, the total-CCC and inter-CCC are defined as

$$\begin{aligned}
 CCC_{total}^R = \rho_c^R &= 1 - \frac{\sum_{j=1}^{J-1} E(Y_{ij} - Y_{iJ})^2}{\sum_{j=1}^{J-1} E_I(Y_{ij} - Y_{iJ})^2} = \frac{2 \sum_{j=1}^{J-1} \sigma_j \sigma_J \rho_{jJ}}{\sum_{j=1}^{J-1} [\sigma_j^2 + \sigma_J^2 + (\mu_j - \mu_J)^2]} \\
 &= \frac{2 \sum_{j=1}^{J-1} \sigma_{Bj} \sigma_{BJ} \rho_{\mu jJ}}{\sum_{j=1}^{J-1} [2\sigma_{Bj} \sigma_{BJ} + (\mu_j - \mu_J)^2 + (\sigma_{Wj} - \sigma_{WJ})^2 + \sigma_{Wj}^2 + \sigma_{WJ}^2]}; \\
 CCC_{inter}^R = \rho_c(\mu)^R &= 1 - \frac{\sum_{j=1}^{J-1} E(\mu_{ij} - \mu_{iJ})^2}{\sum_{j=1}^{J-1} E_I(\mu_{ij} - \mu_{iJ})^2} \\
 &= \frac{2 \sum_{j=1}^{J-1} \sigma_{Bj} \sigma_{BJ} \rho_{\mu jJ}}{\sum_{j=1}^{J-1} [2\sigma_{Bj} \sigma_{BJ} + (\mu_j - \mu_J)^2 + (\sigma_{Bj} - \sigma_{BJ})^2]}
 \end{aligned}$$

They are related via

$$\frac{1}{\rho_c^R} = \frac{1}{\rho_c(\mu)^R} + \frac{1}{\gamma^{R*}}$$

where

$$\frac{1}{\gamma^{R*}} = \frac{\sum_{j=1}^{J-1} (\sigma_{Wj}^2 + \sigma_{WJ}^2)}{2 \sum_{j=1}^{J-1} \sigma_{Bj} \sigma_{BJ} \rho_{\mu jJ}} = \sum_{j=1}^J \omega_j^R \frac{1 - \rho_j^I}{\rho_j^I},$$

with $\omega_j^R = \sigma_{Bj}^2 / (2 \sum_{j=1}^{J-1} \sigma_{Bj} \sigma_{BJ} \rho_{\mu jJ})$, $j = 1, \dots, J-1$ and $\omega_J^R = (J-1) \sigma_{BJ}^2 / (2 \sum_{j=1}^{J-1} \sigma_{Bj} \sigma_{BJ} \rho_{\mu jJ})$ is the weighted sum of the odds of $1 - \rho_j^I$. We note that for $J = 2$, the CCCs for the case with a reference observer are the same as for the ones without reference observer.

We can interpret the CCC_{total} as a measure of overall agreement and the CCC_{total}^R as a measure of validity. The CCC_{inter} assesses systematic shifts and $CCC_{j,intra}$ assesses the precision of the j th observer. All of these indices are scaled relative to the between-subject variability (σ_{Bj}^2) and would produce a high value for a population with large between-subject variability.

For the special case of $J = 2$ and $K = 1$, Lin (1989) defined $\rho_{12} = \text{corr}(Y_{i1}, Y_{i2})$ and $\chi_a = 2\sigma_1\sigma_2 / (2\sigma_1\sigma_2 + (\mu_1 - \mu_2)^2 + (\sigma_1 - \sigma_2)^2)$ as the precision and accuracy components of the CCC, respectively. These two components are scaled indices of systematic bias and precision. We note that $\rho_{12} = \rho_{\mu 12} \sqrt{\rho_1^I \rho_2^I}$ and thus if $\rho_{\mu 12} = 1$, $\rho_{1,2}$ is an inverse of intra-CCCs that are scaled indices of within-subject variabilities σ_{Wj}^2 . We note that χ_a assesses the systematic bias due to both location and scale shifts.

If the agreement between multiple observers is not satisfactory based on the CCCs, it is useful to compare the pairwise CCCs of multiple observers to find which observers do not agree well. One may be also interested in whether there is a trend in the agreement, or whether the observers have good or poor agreement in particular subgroups. Barnhart and Williamson (2001) used the GEE approach to compare these pairwise CCCs and to investigate the impact of covariates. One should be cautious in interpreting CCCs when covariates are used in modeling. Because CCCs depend on between-subject variability, one must ensure that between-subject variability is similar across the range of covariate values. CCC values may

decrease due to decreased between-subject variability when considering subgroups of different covariate values. The GEE approach can be extended to compare CCCs over time in cases where time is a covariate. Specifically, for data with repeated measures, and using $J = 2$ as an example, CCCs can be computed at each time point, and one can model CCCs with Fisher's Z-transformation as a function of time. If there is no time effect, one can obtain a common CCC that is aggregated over time. This common CCC is similar to the straight average of CCCs computed from each time point, but differs from the CCC discussed below by Chinchilli et al. (1996) and King et al. (2007a) for repeated measures.

For two observers ($J = 2$), Chinchilli et al. (1996) proposed weighted CCC, $CCC(\omega)$, and King et al. (2007a) proposed repeated measures CCC, $CCC(rm)$, for data with repeated measures. The repeated measurements can occur due to multiple visits in longitudinal studies, or arise when multiple subsamples are split from the original sample. The repeated measures are not considered replications because they may not be independently and identically distributed, given subject or sample. Suppose that for each observer, there are p repeated measures for a subject, rather than K replications. There are then a total of p^2 pairs of measurements between the two observers. Intuitively, one can assess agreement by using an agreement matrix $CCC_{p \times p}$ consisting of all pairwise CCCs based on the p^2 pairs of measurements using Lin's (1989) original definition of CCC. These pairwise CCCs are concerned with agreement between observations of the two observers who made measurements at the same or at different time points, although the agreement between measurements at the same time is probably the most interesting. One may be interested in assessing agreement between measurements at different time points if the true value of the subject is constant at a prespecified length of time when the observations are taken. The question is how to aggregate the information by using one summary index, instead of these p^2 numbers.

Chinchilli et al. (1996) constructed the $CCC(\omega)$ as a weighted average of q CCCs where the q CCCs are defined as the CCCs between two observers based on q predicted (transformed) new variables obtained from a random-coefficient generalized multivariate ANOVA model. The model-based transformation of observations is most useful if the observers take different numbers of repeated measures over time or across subjects. In the case where each observer takes p measurements at the same p time points, one would not need the transformation (or the transformation is identity) and we have $q = p$. The weights used in Chinchilli et al. (1996) are the inverses of within-subject variabilities. If the within-subject variabilities are the same for all subjects, then the $CCC(\omega)$ is the same as the straight average of the p pairwise CCCs between two observers who made measurements at one of the p time points. In this special case, the $CCC(\omega)$ is similar to the common CCC obtained by using the GEE approach (Barnhart and Williamson, 2001) described above in cases where there is no time effect.

Let $\mathbf{Y}_{i1} = (Y_{i11}, \dots, Y_{i1p})'$ and $\mathbf{Y}_{i2} = (Y_{i21}, \dots, Y_{i2p})'$ be the observations made by two observers, respectively. King et al. (2007a) defined repeated measures CCC by using a distance matrix \mathbf{D} as

$$CCC_{\mathbf{D}} = \rho_{c,rm} = 1 - \frac{E[(\mathbf{Y}_{i1} - \mathbf{Y}_{i2})' \mathbf{D} (\mathbf{Y}_{i1} - \mathbf{Y}_{i2})]}{E_I[(\mathbf{Y}_{i1} - \mathbf{Y}_{i2})' \mathbf{D} (\mathbf{Y}_{i1} - \mathbf{Y}_{i2})]}.$$

\mathbf{D} can be thought of as a weighting matrix. Rather than taking the weighted average of the pairwise CCCs, the repeated measure of CCC here is the ratio of the weighted numerator and weighted denominator based on the pairwise CCCs. This weighting approach is probably more stable than the weighted average of the resulting divisions of numerators and denominators from the pairwise CCCs. Nevertheless, the repeated measures CCC can probably be rewritten as a weighted average of the pairwise CCCs, where the weights would be a function of \mathbf{D} and population parameters. Four options for the \mathbf{D} matrix are considered in King et al. (2007a) with the two most interesting cases of (1) identity matrix $\mathbf{D} = \mathbf{I}_{p \times p}$ and (2) matrix with all entries of one, $\mathbf{D} = (d_{kk'})$ with $d_{kk'} = 1$. The first option gives the same weight to observations taken at the same time and zero weight to all others; the second option gives the same weight to observations taken at the same time and at different times.

King et al. (2007b) extended the repeated measures CCC to a class of repeated measures of agreement as

$$CCC_{\mathbf{D},\delta} = \rho_{c,rm}(\delta) = 1 - \frac{E[\sum_{k=1}^p \sum_{k'=1}^p d_{kk'} |Y_{i1k} - Y_{i2k}|^\delta |Y_{i1k'} - Y_{i2k'}|^\delta]}{E_I[\sum_{k=1}^p \sum_{k'=1}^p d_{kk'} |Y_{i1k} - Y_{i2k}|^\delta |Y_{i1k'} - Y_{i2k'}|^\delta]},$$

where it reduces to the repeated measures CCC if $\delta = 1$ and to a parameter comparable to repeated measures version of kappa (Cohen, 1960, 1968) index for assessing agreement between categorical measurements if $\delta = 0$. It would be of interest to extend the repeated measures CCC to the general case of J observers. King et al. (2007a,b) did not consider agreement between observations made by the same observer at different time points. With an additional assumption that takes time into account, one may be able to define and CCC_{intra} and CCC_{inter} for repeated measures and to derive the relationship of CCC_D with CCC_{intra} and CCC_{inter} , when D is the identity matrix. It would also be of interest to extend the repeated measures CCC to the case with one observer as a reference.

Estimation of CCCs can be done by plugging in the sample counterparts of the population parameters. These estimates can also be obtained from SAS procedure MIXED (Barnhart et al., 2005a; Carrasco and Jover, 2003a) by using the following sample codes:

```

/* * if K = 1 */
proc mixed;
  model Y = observer/s;
  random id;
  run;
/* * if K > 1 */
proc mixed;
  class id observer;
  model Y = observer/noint s;
  random observer/G subject = id type = un V;
  repeated/R group = observer;
  run;

```

where the solution in the model statement provides the estimates for μ_j 's, the G matrix provides the estimates for σ_{Bj}^2 and $\rho_{u_{jj'}}$, and the R matrix provides the estimates for $\sigma_{w_{jj'}}^2$. Carrasco and Jover (2003a) noted that $(Y_{\bullet,j\bullet} - Y_{\bullet,j'\bullet})^2$

is a biased estimator of $(\mu_j - \mu_{j'})^2$ because $E(Y_{\bullet j\bullet} - Y_{\bullet j'\bullet})^2 = (\mu_j - \mu_{j'})^2 + \text{Var}(Y_{\bullet j\bullet}) + \text{Var}(Y_{\bullet j'\bullet}) - 2\text{Cov}(Y_{\bullet j\bullet}, Y_{\bullet j'\bullet}) = (\mu_j - \mu_{j'})^2 + \sigma_{Bj}^2/n + \sigma_{Wj}^2/(nK) + \sigma_{Bj'}^2/n + \sigma_{Wj'}^2/(nK) - 2\sigma_{Bj}\sigma_{Bj'}\rho_{\mu jj'}/n$. The bias may be negligible for moderate-to-large sample sizes.

Parametric, semiparametric, and nonparametric approaches have been proposed for inference of CCCs. If we assume normality of the data, asymptotic distribution of the estimated CCCs may be used for inference (Carrasco and Jover, 2003a; Lin, 1989). The semiparametric approach is available via GEE approach (Barnhart and Williamson, 2001; Lin et al., 2007) and the nonparametric approach was proposed by King and Chinchilli (2001a) using U-statistics. Williamson et al. (2007) proposed and compared permutation and bootstrap tests for testing equalities of CCCs under various conditions. The permutation test is only valid if the joint distributions of the observations under these various conditions are the same, an assumption that may be difficult to verify. Thus, bootstrap-based test may be preferred.

3.3.2.1. Comparison of CCC and ICC. While (total) CCC and ICC are similar indices, there are some differences between them: (1) the ICC has been proposed for both fixed and random observers, while the CCC usually treats the observers fixed; and (2) the ICC requires ANOVA model assumptions, while the CCC does not. However, in specific cases, the ICC and CCC are the same or have similar values (Carrasco and Jover, 2003a; Nickerson, 1997). For example, if there are no replications, Carrasco and Jover (2003a) demonstrated that ICC_2 is the same as total CCC even without the ANOVA model assumption. In general, if the ANOVA model assumptions are correct, the CCC under this model reduces to the ICC defined by this ANOVA model.

ICC estimators are based on unbiased estimates for the parameters used in the assumed ANOVA models. However, it is not clear what the ICC estimators are estimating if the assumed ANOVA models are not correct. Using a general model of $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$, Chen and Barnhart (2007) compared expected values of the three ICC estimators in Section 3.3.1 under this general model to the total CCC under the same model. They approximated the expected values of the ICC estimators by taking the expectation of the numerator and denominator under this general model. They found that if there are no replications ($K = 1$), $E(\widehat{ICC}_1)$ may be smaller or larger than the CCC, where for $J = 2$,

$$E(\widehat{ICC}_1) = \frac{2\rho\sigma_1\sigma_2 - 0.5(\mu_1 - \mu_2)^2}{\sigma_1^2 + \sigma_2^2 + 0.5(\mu_1 - \mu_2)^2}.$$

We note that this $E(\widehat{ICC}_1)$ is the same as the random marginal agreement coefficient (RMAC) proposed by Fay (2005) for continuous response. For ICC_2 and ICC_3 , we have equality, i.e., $E(\widehat{ICC}_2) = E(\widehat{ICC}_3) = CCC$. For data with replications ($K > 1$), Chen and Barnhart (2007) found that under the general model, $E(\widehat{ICC}_3) = CCC$. However, the expected ICCs for the first two ICC estimators depend on the number of replications, K , whereas the CCC does not depend on K . For the special case with homogeneous between-and within-subject variability; i.e., $\sigma_{Bj}^2 = \sigma_B^2$, $\sigma_{Wj}^2 = \sigma_W^2$ both $E(\widehat{ICC}_1)$ and $E(\widehat{ICC}_2)$ are increasing functions of K . $E(\widehat{ICC}_1)$ starts with a value that may be less or greater than the CCC at $K = 1$ and increases quickly to the limit

(as $K \rightarrow \infty$) that exceeds the CCC. $E(\widehat{ICC}_2)$ equals the CCC at $K = 1$ and increases quickly to the limit (as $K \rightarrow \infty$) that also exceeds the CCC.

Note that even in the case of $E(\widehat{ICC}) = CCC$ under the general model, we may still have $\widehat{CCC} \leq \widehat{ICC}$, because the plug-in estimator of CCC, \widehat{CCC} , is biased due to a biased estimation of $(\mu_j - \mu_{j'})^2$ via $(Y_{\bullet j\bullet} - Y_{\bullet j'\bullet})^2$ (Carrasco and Jover, 2003a; Nickerson, 1997). We have $\widehat{CCC} = \widehat{ICC}$ only if a bias correction term is used for estimating $(\mu_j - \mu_{j'})^2$ in the CCC.

3.3.3. Coefficient of individual agreement. The CCC is known to depend on between-subject variability that may result from the fact that it is scaled relative to the maximum disagreement defined as the expected squared difference under independence. Barnhart and colleagues (Barnhart et al., 2007a; Haber and Barnhart, 2007) began looking for a scaled agreement index, the coefficient of individual agreement (CIA), which is scaled relative to the minimum or acceptable disagreement, with the goal of establishing interchangeability of observers. Before considering observers for comparison, one must assume that the replication errors of the observers are acceptable. This is especially true for the reference observer. Thus, they used the disagreement between replicated measurements within an observer as a yardstick for acceptable disagreement. Intuitively, interchangeability is established only if individual measurements from different observers are similar to replicated measurements of the same observer. In other words, the individual difference between measurements from different observers is relatively small, so that this difference is close to the difference of replicated measurements within an observer. This concept of individual agreement is closely linked to the concept of individual bioequivalence in bioequivalence studies (Anderson and Hauck, 1990, Schall and Luus, 1993).

Replicated measurements by observers are needed for the CIAs for the purpose of estimation and inference only. Regardless of the number of replications, CIAs assesses agreement between observers when each observer evaluates each subject only once. The coefficients are intended to be used to decide whether a single measurement made by one observer can be replaced by a single measurement made by another observer in practice, when each subject is evaluated only once by each observer. Let $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$ with the assumptions in Section 3.1. Barnhart et al. (2007a) defined the CIAs for cases of no reference observer and the J th observer as a reference with

$$\begin{aligned}
 CIA^N = \psi^N &= \frac{\sum_{j=1}^J E(Y_{ijk} - Y_{ijk'})^2 / 2}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J E[(Y_{ijk} - Y_{ijk'})^2] / (J-1)} \quad (\text{where } k \neq k') \\
 &= \frac{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J (\sigma_{Wj}^2 + \sigma_{Wj'}^2)}{\sum_{j=1}^{J-1} \sum_{j'=j+1}^J [2(1 - \rho_{\mu jj'}) \sigma_{Bj} \sigma_{Bj'} + (\mu_j - \mu_{j'})^2 + (\sigma_{Bj} - \sigma_{Bj'})^2 + \sigma_{Wj}^2 + \sigma_{Wj'}^2]} \\
 CIA^R = \psi^R &= \frac{E(Y_{ijk} - Y_{ijk'})^2 / 2}{\sum_{j=1}^{J-1} E[(Y_{ijk} - Y_{ijk'})^2] / (J-1)} \quad (\text{where } k \neq k') \\
 &= \frac{\sigma_{WJ}^2}{\sum_{j=1}^{J-1} [2(1 - \rho_{\mu jJ}) \sigma_{Bj} \sigma_{BJ} + (\mu_j - \mu_J)^2 + (\sigma_{Bj} - \sigma_{BJ})^2 + \sigma_{Wj}^2 + \sigma_{WJ}^2]},
 \end{aligned}$$

respectively. These are the CIAs when the MSD function is used as a disagreement function. For $J = 2$, Haber and Barnhart (2007) extended the CIAs to the general disagreement function and illustrated the methodology with various disagreement functions. Barnhart et al. (2007a) showed that there are one-to-one relationships between the CIA^N and the two previously proposed agreement indices by Haber et al. (2005) and Shao and Zhong (2004).

Estimates of CIAs can be obtained by the method of moment and can be computed through several ANOVA models (Barnhart et al., 2007a). Barnhart et al. (2007a) proposed a nonparametric approach for inference and showed that the nonparametric approach is similar to the bootstrap approach. This nonparametric approach is also similar to the U-statistics used by King and Chinchilli (2001a) for the CCCs.

3.3.3.1. Comparison of CIA and CCC. Barnhart et al. (2007b) compared the CCC and the CIA for data with replications when there is no reference observer. To highlight the similarities and differences of the two coefficients, we assume that there are only two observers and that $\sigma_{B1}^2 = \sigma_{B2}^2 = \sigma_B^2$ and $\sigma_{W1}^2 = \sigma_{W2}^2 = \sigma_W^2$. We can write both coefficients in terms of the difference of the means ($\mu_1 - \mu_2$), the between- and within-subjects variances, and the correlation coefficient ($\rho_{\mu 12}$). The total CCC and the CIA for the case of no reference can be written as:

$$\rho_c = \frac{2\sigma_B^2\rho_{\mu 12}}{(\mu_1 - \mu_2)^2 + 2(\sigma_B^2 + \sigma_W^2)}, \quad \psi^N = \frac{2\sigma_W^2}{(\mu_1 - \mu_2)^2 + 2(1 - \rho_{\mu 12})\sigma_B^2 + 2\sigma_W^2}.$$

Hence, both coefficients increase as the correlation increases and decrease as the overall location shift increases. The CCC increases when the between-subjects variability increases and the within-subjects variability decreases. The CIA, on the other hand, increases when the within-subjects variability increases and the between-subjects variability decreases. However, Barnhart and colleagues found that the CIA is less dependent than the CCC on the relative magnitude, σ_B^2/σ_W^2 , of the between- and within-subjects variability. In general, the CCC and the CIA are related as

$$\psi^N = \rho_c / [(1 - \rho_c)\gamma],$$

where $\gamma = 2\sigma_{B1}\sigma_{B2}\rho_{\mu 12}/(\sigma_{W1}^2 + \sigma_{W2}^2)$.

These properties of the CCC and the CIA continue to apply when there are more than two observers, none of whom is considered as a reference. Comparison of a new CCC (where one observer is the reference) to the corresponding CIA (with the same observer as a reference) leads to the same conclusion as the comparison of the ordinary CCC with the CIA when there is no reference observer.

3.3.4. Dependability coefficient and generalizability coefficient. Generalizability theory (GT) extends the concept of reliability based on classical theory (CT) to account for various sources of variability and different kinds of decisions. We first introduce the CT that defines the traditional reliability coefficient, followed by a single-facet design and a multi-facet design in GT, where we discuss the definition of dependability coefficient and generalizability coefficient.

In CT, an observation is decomposed as the sum of the subject's true score, plus the random measurement error. In the simple case that an observation is made

by an observer, let Y_{ij} be the observations made by the j th observer on subject i . Then, the one-way ANOVA model $Y_{ij} = \mu_i + \epsilon_{ij} = \mu + \alpha_i + \epsilon_{ij}$ is used in CT, where observations Y_{ij} are assumed to be parallel (see Section 2.3 under reliability). The reliability is defined as ICC_1 (see Section 3.3.1) and can be viewed either as the ratio of the true score variance ($Var(\mu_i) = \sigma_\alpha^2$) over the observed variance ($Var(Y_{ij}) = \sigma_\alpha^2 + \sigma_\epsilon^2$) or as correlation $Corr(Y_{ij}, Y_{ij'})$.

In GT, the assumption of parallel observations is relaxed by decomposing an observation as sum of subject's universe score and the multiple sources of components that contribute to the variability of the observed value, where the universe score is similar to the concept of true score in CT. The subject is usually called the *facet of differentiation* or the *object (facet) of measurement* and the components are called *facets of generalization*. The levels of facets of generalization are called *conditions*. The *universe* is defined as all observations made under all conditions of the facets of generalization. The *universe score* is the average of all possible (usually infinite) observations in the universe, similar to the true score in CT. The simplest case is the single facet design, where the observer is the only facet of generalization. The universe is all observations made by all (possibly infinite) observers, and the universe score is the average of these observations. GT consists of generalization studies (G-studies) and decision studies (D-studies), where G-studies investigate the variance components due to facets of differentiation and facets of generalization, while D-studies investigate various designs based on particular decisions. In general, one G-study is planned to include all possible facets of generalization (i.e., sources of possible variabilities), while multiple D-studies can be formulated by considering the variance components obtained from a single G-study in various ways.

One key convention in GT is that all facets of generalization are considered as random effects. Thus, the observer is treated as random and the GT extends the reliability concept in Section 3.3.1 for the case of the random observer only. Fixed facets of generalization are discussed in Molenberghs et al. (2007).

Let Y_{ijk} be the k th replication ($k = 1, \dots, K$) made by observer j on subject i . Usually, $K = 1$ in the GT, but we allow general K for comparison with the reliability concept in Section 3.3.1 that includes both $K = 1$ and $K > 1$. In a single-facet study, the G-study may consider the following three random effect models: (1) $Y_{ijk} = \mu + \alpha_i + \epsilon_{ijk}$; (2) $Y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon_{ijk}$; and (3) $Y_{ijk} = \mu + \alpha_i + \beta_j + (\alpha\beta)_{ij} + \epsilon_{ijk}$, where $\mu_i = \mu + \alpha_i$ is the universe score. Model (1) is usually not considered in a G-study because the observer is recognized as a facet and thus should always be considered as a factor in the ANOVA model. In general, the G-study uses model (2) if there are no replications and model (3) if there are replications. We include all three models for comparison with the ICCs. The G-study uses these ANOVA models to estimate and perform inference on variance components of $Var(\mu_i) = \sigma_\alpha^2$, $Var(\beta_j) = \sigma_\beta^2$, $Var(\alpha\beta_{ij}) = \sigma_{\alpha\beta}^2$, $Var(\epsilon_{ij}) = \sigma_\epsilon^2$ due to facet of differentiation (subject) and facets of generalization (observer and replication), respectively.

In D-studies, one must decide (1) whether the decision is absolute or relative and (2) whether a single observation, or average of several observations of a subject, is used in practice. An *absolute decision* is concerned with obtaining the estimate of a subject's universe score regardless of another subject's universe score while *relative decision* is concerned with rank-ordering subjects. Thus, the absolute decision is relevant to absolute agreement considered in this paper, while the relative decision

is relevant to consistency agreement, such as ICC, for consistency. As mentioned in the begining of Section 3.1, we only consider the case that a single observation will be used in practice. Therefore, we focus our attention on the D-study with absolute decision, with the purpose of using a single observation in practice. Coefficients that depend on relative decision with the purpose of using a single observation in practice are mentioned here for contrast. Based on the absolute decision, *dependability coefficient* (DC) is defined as

$$\rho_{DC} = \frac{\text{variance of universe score based on facet of differentiation}}{\text{variance of universe score} + \text{variance of absolute error}}.$$

Based on the relative decision, *generalizability coefficient* (GC) is defined as

$$\rho_{GC} = \frac{\text{variance of universe score based on facet of differentiation}}{\text{variance of universe score} + \text{relative error}}.$$

In the single-facet design, the universe score for subject i is $\mu_i = \mu + \alpha_i$, and the absolute error (Δ_{ijk}) is the difference between the observed score and the universe score, $\Delta_{ijk} = Y_{ijk} - \mu_i$. The relative error (δ_{ijk}) is defined as the difference of the subject's observed deviation score ($Y_{ijk} - E(Y_{ijk} | j, k) = Y_{ijk} - \mu - \beta_j$) and the universe deviation score ($\mu_i - \mu$), thus $\delta_{ijk} = Y_{ijk} - \mu_i - \beta_j$. Intuitively, the relative error does not contain the observer effect, because this effect does not affect the subject's ranking. Table 1 shows the dependability coefficient and generalizability coefficient based on the three different ANOVA models. It can be shown that $\rho_{DC} = \text{corr}(Y_{ijk}, Y_{ij'k'})$ and $\rho_{GC} = \text{corr}(Y_{ijk}, Y_{ij'k'} | j, j', k, k')$, where $j \neq j', k \neq k'$ for all three models (Molenberghs et al., 2007). Thus, the DC can be interpreted as correlation and the GC can be interpreted as conditional correlation.

Molenberghs et al. (2007) also considered the single-facet design with a fixed observer and the facet of generalization as the replication. One may consider separate ANOVA models to obtain DC and GC for each observer; however, if model (3) is used, the DC and GC for each observer is the same and equal to

$$\rho_{\text{test-retest}} = \text{corr}(Y_{ijk}, Y_{ijk'} | j) = \text{corr}(Y_{ijk}, Y_{ijk'} | j, k, k') = \frac{\sigma_{\alpha}^2 + \sigma_{(\alpha\beta)}^2}{\sigma_{\alpha}^2 + \sigma_{(\alpha\beta)}^2 + \sigma_{\epsilon}^2}$$

that is considered as test-retest reliability, which is the same as the intra-CCC by Lin et al. (2007).

Table 1 Dependability and Generalizability Coefficients for one-facet design

G-study models	D-study coefficients		
	ρ_{DC}	ρ_{GC}	Relation with ICC
(1) $Y_{ijk} = \mu + \alpha_i + \epsilon_{ijk}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\epsilon}^2}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\epsilon}^2}$	$\rho_{DC} = \rho_{GC} = ICC_1$
(2) $Y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon_{ijk}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\beta}^2 + \sigma_{\epsilon}^2}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\epsilon}^2}$	$\rho_{DC} = ICC_2, \rho_{GC} = ICC_1$
(3) $Y_{ijk} = \mu + \alpha_i + \beta_j + \alpha\beta_{ij} + \epsilon_{ijk}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\beta}^2 + \sigma_{\alpha\beta}^2 + \sigma_{\epsilon}^2}$	$\frac{\sigma_{\alpha}^2}{\sigma_{\alpha}^2 + \sigma_{\alpha\beta}^2 + \sigma_{\epsilon}^2}$	$\rho_{DC} = ICC_3, \rho_{GC} = ICC_3c$

It is now easy to extend the single-facet design to the multifacet design. For illustration, let Y_{ijtc} denote the observation made by observer j at time t on subject i , with covariate value c . The facets of generalization are observer, time, and covariate. The G-study decomposes Y_{ijtc} as the sum of various sources of variability with model (4) below:

$$(4) \quad Y_{ijtc} = \mu + \alpha_i + \beta_j + \tau_t + \theta_c + (\alpha\beta)_{ij} + (\alpha\tau)_{it} + (\alpha\theta)_{ic} \\ + (\beta\tau)_{jt} + (\beta\theta)_{jc} + (\tau\theta)_{tc} + (\alpha\beta\tau)_{ijt} + (\alpha\beta\theta)_{ijc} + (\alpha\tau\theta)_{itc} + (\beta\tau\theta)_{jtc} + \epsilon_{ijtc}$$

The DC and GC from the D-studies with absolute and relative errors are

$$\rho_{DC} = \frac{\sigma_\alpha^2}{\sigma_\alpha^2 + \sigma_\beta^2 + \sigma_\tau^2 + \sigma_\theta^2 + \sigma_{(\alpha\beta)}^2 + \sigma_{(\beta\theta)}^2 + \sigma_{(\tau\theta)}^2 + \sigma_{(\alpha\tau\theta)^2} + \sigma_{(\beta\tau\theta)}^2 + \sigma_\epsilon^2}$$

$$\rho_{GC} = \frac{\sigma_\alpha^2}{\sigma_\alpha^2 + \sigma_{(\alpha\beta)}^2 + \sigma_{(\alpha\tau\theta)}^2 + \sigma_\epsilon^2}.$$

respectively. One should note that these coefficients now assess interchangeability for the measurements by different observers at different time points and different covariate conditions, not only for interchangeability of the observers.

Other DCs or GCs may be formed if we consider different facets of differentiation other than subject; e.g., subject-by-country, or subject-by-observer. Vangeneugden et al. (2005) and Molenberghs et al. (2007) extended the concepts of ICC and GT to longitudinal data in the framework of a linear mixed-effect model with and without serial correlation. The mixed effect model allows for adjustment of fixed effects from covariates such as treatment. In their framework, both the ICC and DC are expressed as correlations, while the GC is expressed as a conditional correlation. In the simple case where Y_{ijt} is the observation made by observer j on subject i at time t , the mixed model is $Y_{ijt} = \mu + \alpha_i + \beta_j + (\alpha\beta)_{ij} + \omega_{it} + \epsilon_{ijt}$, where ω_{it} accounts for serial correlation. They first defined ICC as $ICC = corr(Y_{ijt}, Y_{ijt'})$ for test-retest reliability and then defined three DCs and three GCs for overall, test-retest, and inter-rater respectively as

$$\rho_{DC} = corr(Y_{ijt}, Y_{ijt'}), \quad \rho_{GC} = corr(Y_{ijt}, Y_{ijt'} | j, j', t, t')$$

$$\rho_{DC, test-retest} = corr(Y_{ijt}, Y_{ijt'}), \quad \rho_{GC, test-retest} = corr(Y_{ijt}, Y_{ijt'} | j, t, t')$$

$$\rho_{DC, inter-rater} = corr(Y_{ijt}, Y_{ijt'}), \quad \rho_{GC, inter-rater} = corr(Y_{ijt}, Y_{ijt'} | j, j', t).$$

If the repeated measurements are treated as replicated measurements by setting $\omega_{it} = 0$ in the model, then $\rho_{DC} = \rho_{DC, inter-rater}$. Furthermore, ρ_{DC} and $\rho_{DC, test-retest}$ correspond to the total CCC and intra-CCC, respectively, if we set $\sigma_\beta^2 = \sum_{j=1}^J (\mu_j - \mu_\bullet)^2 / (J - 1)$. For data with repeated measures, $\rho_{DC, inter-rater}$ has an interpretation similar to the CCC_D by King et al. (2007a) with corresponding D matrix. We should point out that the models used in GT assume the same between-subject variability, while the CCC allows separate between-subject variability for each observer.

In summary, GT provides a flexible way to investigate interchangeability of levels in more than one factor (such as observer). However, the DC shares the

same property as the ICC and CCC, in that its value increases as the between-subject variability increases. Thus, one cannot compare the coefficients formed from different G-studies where the populations differ greatly.

3.3.5. Comments. In summary, the scaled agreement indices of ICC, CCC, CIA, and DC are all standardized to have values between -1 and 1 . The ICC, CCC, and DC are related and depend on between-subject variability and may produce high values for heterogeneous populations (Atkinson and Nevill, 1997; Bland and Altman, 1990). The ICC can accommodate multiple fixed and random observers, but is not suited for cases with a reference observer or for data with repeated measures, without additional assumptions. The CCC is mainly developed for fixed observers and reduces to the ICC for fixed observers under additional assumptions. The CCC formulation may be used for the case with random observers, but additional care is needed for inference. The DC is an extension of the ICC for random observers that can be used for situations with multiple factors, and for data with repeated measurements. There has been limited development of the DC for fixed observers; nothing has been developed for the case with a reference observer. The CIA is fundamentally different from the ICC, CCC, and DC because it uses within-subject, rather than between-subject, variability as the scaling factor. It is possible to have high ICC/CCC/DC value and low CIA value (and vice versa) from the same data set. See Barnhart et al. (2007b) for additional details.

4. DISCUSSION

We have reviewed the concepts and statistical approaches for assessing agreement with continuous measurements based on classification of (1) descriptive tools, (2) unscaled indices, and (3) scaled indices. An alternative classification used in individual bioequivalence literature (Chen, 1997) also may be considered: (1) aggregated criteria, (2) disaggregated criteria, and (3) probability criteria. The classification of aggregated criteria is based on an index that combines different sources of possible disagreement among observers. Sources of disagreement may arise from differing population means, differing between-subject variances, differing within-subject variances among observers, poor correlation between measurements made by observers, and large subject-by-observer interaction. Disaggregated criteria examine these various possible sources of disagreement separately (see Lee et al., 1989). Except for the CP and TDI, which are based on probability criteria, all other indices reviewed here are aggregated indices. Our descriptive tools are intuitive approaches to disaggregated criteria. As pointed out by Chen (1997), aggregated criteria have the advantage of balancing different sources of disagreement while disaggregated criteria may have the advantage of identifying the actual source of disagreement if the agreement is not satisfactory. However, disaggregated criteria may encounter the problem of multiple testing if the criteria do not use one procedure to test multiple hypotheses together. We did not review disaggregated criteria for assessing agreement that takes into account multiple testing; readers are instead directed to the literature that were based on the intersection-union principle (Carrasco and Jover, 2003b, 2005a; Choudhary and Nagaraja, 2005a) or other single procedures (Bartko, 1994).

We summarize here the unscaled (Table 2) and scaled (Table 3) agreement indices according to whether there is an existing method for different types of data structure. The tables indicate that future research is needed for agreement indices where there are repeated measures, covariates, and the existence of a reference observer. Furthermore, most methodologies were developed for fixed observers; thus, further research is needed for the case of random observers. The case of random observers should be considered if different observers from a pool of observers, rather than the same limited observers, are used in practice or in research. Recently, Barnhart et al. (2005b) used the CCC index and Choudhary (2007a) used the CP and TDI indices to assess agreement of measurements subject to censoring due to the limit of detection. Ying and Manatunga (2007) and Liu et al. (2005) used the CCC index to assess agreement of time measurements that are subject to censoring, with nonparametric and parametric estimation approaches, respectively. Clearly, further research is needed to understand assessing agreement for censored data. King and Chinchilli (2001b) proposed robust estimators of CCC to reduce the influence of outliers and Carrasco et al. (2007) evaluated performance of various estimators of CCC with skewed data. More research is also needed for assessing agreement with data containing outliers.

Unscaled indices have the advantage of interpretation based on the original unit, but it may prove difficult to ascertain the limit for acceptable agreement without sufficient knowledge of the measurement variable and measurement unit. Scaled indices have the advantage of judging the degree of agreement based on standardized value, but the agreement values may not be compared across very different populations, and sometime artificially high or low agreement values may be obtained due to the dependence of these indices (except the CIA) on between-subject variability. If there are only two observers ($J = 2$), we note that there is

Table 2 Existing methods (Yes, No) for unscaled agreement indices comparing between or within J observers under different data structures

Index	Data structure						Covariates
	No replications		Replications		Repeated measures		
	$J = 2$	$J > 2$	$J = 2$	$J > 2$	$J = 2$	$J > 2$	
No reference observer							
MSD	Yes	Yes	Yes	Yes	No	No	No
Repeatability	NA	NA	Yes	Yes	Yes	No	Yes
Reproducibility	Yes	Yes	Yes	Yes	No	No	No
LOA	Yes	No	Yes	No	Yes	No	Yes
CP	Yes	Yes	Yes	Yes	Yes	No	No
TDI	Yes	Yes	Yes	Yes	Yes	No	Yes
With reference observer							
MSD	Yes	No	Yes	No	No	No	No
Repeatability	NA	NA	Yes	Yes	No	No	No
Reproducibility	Yes	No	Yes	No	No	No	No
LOA	Yes	No	Yes	No	Yes	No	No
CP	Yes	No	Yes	No	No	No	No
TDI	Yes	No	Yes	No	No	No	No

Table 3 Existing methods (Yes, No) for scaled agreement indices comparing between or within J observers under different data structures

Index	Data structure						Covariates
	No Replications		Replications		Repeated measures		
	$J = 2$	$J > 2$	$J = 2$	$J > 2$	$J = 2$	$J > 2$	
No reference observer							
ICC	Yes	Yes	Yes	Yes	Yes	Yes	No
CCC	Yes	Yes	Yes	Yes	Yes	No	Yes
CIA	NA	NA	Yes	Yes	No	No	No
DC	Yes	Yes	Yes	Yes	Yes	Yes	No
With reference observer							
ICC	No	No	No	No	No	No	No
CCC	Yes	Yes	Yes	Yes	No	No	No
CIA	NA	NA	Yes	Yes	No	No	No
DC	No	No	No	No	No	No	No

no difference between cases with and without reference observers for most indices, except for CIA (which may be the strength of this index).

If there is a reference observer, agreement indices assess validity. Otherwise, they assess agreement only. It is thus possible that there may be high agreement between observers, but they may agree to the wrong quantity when there is no reference observer or observed true value for comparison. If the focus is not on agreement, but on the relationship between the observed measurements with the true value, the reader is directed to the literature by Lin et al. (2002), where the true value is observed without random error, and to literature on statistical evaluation of measurement errors in method comparison studies (see references in Dunn, 2004) where the true value is observed with random error. Often, the investigation of such a relationship involves assumption of linearity (e.g., in a calibration problem) and/or existence of an instrumental variable (Dunn, 2007; Dunn and Roberts, 1999) that may be difficult to verify in practice. If we believe the assumption of such a linear relationship, one may use disaggregated criteria to test whether there is satisfactory agreement between observers. This approach for assessing agreement may be based on structural equations (latent variables) models (Carrasco and Jover, 2003b, 2005a; Dunn, 1989, 2004; Hawkins, 2002; Kelly, 1985) or a confirmatory factor model (Dunn, 1989, 2004).

For example, under the general model of $Y_{ijk} = \mu_{ij} + \epsilon_{ijk}$ with the J th observer as the reference observer, it is often assumed that $\mu_{ij} = \alpha + \beta\mu_{iJ}$ in the structural equations approach. This implies assumptions of $\mu_j = \alpha + \beta\mu_J$ and $\sigma_{Bj}^2 = \beta^2\sigma_{BJ}^2$. In general, $\sigma_{Bj}^2 = \beta^2\sigma_{BJ}^2$ is usually not required in addition to $\mu_j = \alpha + \beta\mu_J$ in the aggregated approach for assessing agreement. Another similar approach can be found in St. Laurent (1998) and Harris et al. (2001). For illustration, let the J th observer be a reference or gold standard with measurement error. St. Laurent (1998) used the model, $Y_{ij} = Y_{iJ} + \epsilon_{ij}$, to construct an agreement index defined as $\rho_{j,g} = \sqrt{\text{corr}(Y_{ij}, Y_{iJ})} = \text{Var}(Y_{iJ}) / (\text{Var}(Y_{iJ}) + \text{Var}(\epsilon_{ij}))$, where g stands for the gold standard method. Under this model, it can be shown that the pairwise CCC is

$\rho_{c,ij} = 2\text{Var}(Y_{ij}) / (2\text{Var}(Y_{ij}) + \text{Var}(\epsilon_{ij})) = 2\rho_{jg} / (2\rho_{jg} + 1) \geq \rho_{jg}$. The formulation of ρ_{jg} is similar to ICC_1 by treating Y_{ij} as μ_i . If we accept the additive assumption in the model $Y_{ij} = Y_{ij} + \epsilon_{ij}$, the correlation of the observations between the new observer and the reference observer may be used as an alternative agreement index that extends ICC/CCC/DC to the case with a reference observer. If there are multiple observers with one reference observer, one may be also interested in selecting the best observer as compared to the reference observer (Choudhary and Nagaraja, 2005b,c; Hutson et al., 1998; St. Laurent, 1998). If there are $J > 2$ observers and there is no reference, one may be also interested in looking at pairwise agreement between any two observers, especially if the overall agreement is not satisfactory. Furthermore, one may be interested in selecting subgroup of observers who agree well with each other.

Other scaled agreement indices include the within-subject coefficient of variation (Hui and Shih, 1996) and δ coefficients proposed by Zhong and Shao (2003) and Shao and Zhong (2004). These indices are not scaled to be between -1 and 1 . The within-subject coefficient of variation is repeatability scaled by the mean, rather than between-subject variability. Shao and Zhong (2004) showed that the two δ s are related and the δ coefficient by Shao and Zhong (2004) has a one-to-one correspondence with the CIA index as shown by Barnhart et al. (2007a). As mentioned in Section 3.3.2, the RMAC proposed by Fay (2005) is closely related to $E(\widehat{ICC}_1)$ under the general model. There is also a different version of CCC (Liao, 2003, 2005) based on the concept of area rather than distance.

One implicit assumption among existing methods is that the between- and within-subject variabilities are reasonably stable across the range of measurements. It may be of interest to examine the index's dependency on the magnitude of the true value. Without observed true value, one may examine the indices' dependency on the average magnitude of the observed measurements from the same subject (see Bland and Altman plot, Bland and Altman, 1999).

Sometimes measurement methods or tools are developed to measure more than one variable; e.g., systolic and diastolic blood pressure, multiple characteristics of psychology and psychiatry profiles, multiple outputs of imaging scans, etc. There is a multivariate generalizability theory for multivariate data, where DC or GC indices are aggregated over multiple outcome variables (Brennan, 2001). However, the research on agreement for multivariate data is very limited (see Konishi et al., 1991 for multivariate ICC in genetic study, and Janson and Olsson (2001, 2004) for multivariate CCC in the context of education and psychology). There is a great need for further development in this area.

Sample size calculations for agreement studies have been proposed for indices of ICC (Donner, 1998; Shoukri et al., 2004), CCC (Lin, 1992; Lin et al., 2002), LOA (Lin et al., 1998), and CP or TDI (Choudhary and Nagaraja, 2007; Lin et al., 2007). Further research is needed in the design of agreement studies.

Disagreement between observers will have an effect on the design of clinical trials if the measurements of the observers are used as outcomes. Fleiss (1986) examined the inflation of sample size for a two-arm parallel clinical trial if there is only random measurement error (imprecision), as assessed by ICC_1 . Further research is needed to examine the impact of disagreement on the design of clinical trials where there are both systematic and random measurement errors in the outcome measurements.

In this paper, we have reviewed only the indices for assessing agreement with continuous measurements. Parallel indices were developed for assessing agreement with categorical measurements. For example, Cohen's kappa for binary data and weighted kappa for ordinal data correspond to the CCC for continuous data, and intraclass kappa for ordinal data corresponds to the CCC and ICC_1 (Fleiss and Cohen, 1973; Krippendorff, 1970; Robieson, 1999). Carrasco and Jover (2005b) developed the CCC for count data through the ICC from a generalized linear mixed model. Lin et al. (2007) provided a unified approach to defining total CCC, inter-CCC, and intra-CCC for continuous, binary, or ordinal data. King and Chinchilli (2001a, 2007b) proposed a class of CCC index for continuous or categorical data, including repeated measures. There is also a CIA index, which has been developed for binary data (Haber et al., 2007). Raghavachari (2004) proposed a new measure of agreement for assessing agreement in ratings and rank-order data similar to Kendall's (1948) measure of concordance for ranked-order data. Molenberghs et al. (2007) used generalized linear mixed models to define reliability and generalizability for both continuous and categorical data. We will review the agreement indices for categorical data in the future.

In summary, future research on assessing agreement should be focused on

- Indices for data with repeated measurements, censoring, outliers, and covariates
- Indices for the case of random observers
- Indices for the case with existence of reference
- Investigation of indices' dependency on the range of measurements
- Indices for multivariate data
- Sample size calculation for design of agreement study
- Impact of disagreement on design of clinical trials

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REFERENCES

- American Educational Research Association (AERA), American Psychological Association (APA), National Council on Measurement in Education (NCME). (1999). *The Standards for Educational and Psychological Testing*. Washington, D.C.: American Psychological Association.
- Anderson, S., Hauck, W. W. (1990). Consideration of individual bioequivalence. *Journal of Pharmacokinetics and Biopharmaceutics* 18:259–273.
- Atkinson, G., Nevill, A. (1997). Comment on the use of concordance correlation to assess the agreement between two variables. *Biometrics* 53:775–777.
- Barnhart, H. X., Williamson, J. M. (2001). Modeling concordance correlation via GEE to evaluate reproducibility. *Biometrics* 57:931–940.
- Barnhart, H. X., Haber, M., Song, J. (2002). Overall concordance correlation coefficient for evaluating agreement among multiple observers. *Biometrics* 58:1020–1027.

- Barnhart, H. X., Song, J., Haber, M. (2005a). Assessing intra, inter, and total agreement with replicated measurements. *Statistics in Medicine* 24:1371–1384.
- Barnhart, H. X., Song, J., Lyles, R. (2005b). Assay validation for left censored data. *Statistics in Medicine* 24:3347–3360.
- Barnhart, H. X., Haber, M., Kosinski, A. S. (2007a). Assessing individual agreement. *Journal of Biopharmaceutical Statistics* 17(4):697–719.
- Barnhart, H. X., Haber, M., Lokhnygina, Y., Kosinski, A. S. (2007b). Comparison of concordance correlation coefficient and coefficient of individual agreement in assessing agreement. *Journal of Biopharmaceutical Statistics* 17(4):721–738.
- Bartko, J. J. (1966). The intraclass correlation coefficient as a measure of reliability. *Psychological Reports* 19:3–11.
- Bartko, J. J. (1974). Corrective note to the intraclass correlation coefficient as a measure of reliability. *Psychological Reports* 34:418.
- Bland, J. M., Altman, D. G. (1986). Statistical methods for assessing agreement between two methods of clinical measurement. *Lancet* i:307–310.
- Bland, J. M., Altman, D. G. (1990). A note on the use of the intraclass correlation coefficient in the evaluation of agreement between two methods of measurement. *Computers in Biology and Medicine* 20:337–340.
- Bland, J. M., Altman, D. G. (1992). This week's citation classic: comparing methods of clinical measurement. *Current Contents CM* 20(40) Oct 5:8.
- Bland, J. M., Altman, D. G. (1995). Comparing methods of measurement: why plotting difference against standard method is misleading. *Lancet* 346:1085–1087.
- Bland, J. M., Altman, D. G. (1999). Measuring agreement in method comparison studies. *Statistical Methods in Medical Research* 8:135–160.
- Bland, J. M., Altman, D. G. (2007). Agreement between methods of measurement with multiple observations per individual. *Journal of Biopharmaceutical Statistics* 17(4):571–582.
- Brennan, R. L. (2001). *Generalizability Theory*. New York: Springer.
- Carrasco, J. L., Jover, L. (2003a). Estimating the generalized concordance correlation coefficient through variance components. *Biometrics* 59:849–858.
- Carrasco, J. L., Jover, L. (2003b). Assessing individual bioequivalence using the structural equation model. *Statistics in Medicine* 22:901–912.
- Carrasco, J. L., Jover, L. (2005a). The structural error-in-equation model to evaluate individual bioequivalence. *Biometrical Journal* 47:623–634.
- Carrasco, J. L., Jover, L. (2005b). Concordance correlation coefficient applied to discrete data. *Statistics in Medicine* 24:4021–4034.
- Carrasco, J. L., King, T., Chinchilli, V., Jover, L. (2007). Comparison of concordance correlation coefficient estimating approaches with skewed data. *Journal of Biopharmaceutical Statistics* 17(4):673–684.
- Chen, M. L. (1997). Individual bioequivalence—A regulatory update. *Journal of Biopharmaceutical Statistics* 7:5–11.
- Chen, C., Barnhart, H. X. (2007). Comparison of ICC and CCC for assessing agreement for data without and with replications. ENAR 2007 conference presentation in Atlanta, GA.
- Chinchilli, V. M., Martel, J. K., Kumanyika, S., Lloyd, T. (1996). A weighted concordance correlation coefficient for repeated measures designs. *Biometrics* 52:341–353.
- Choudhary, P. K. (2007a). A tolerance interval approach for assessment of agreement in method comparison studies with repeated measurements. *Journal of Statistical Planning and Inference* DOI: 10.1016/j.jspi.2007.03.056
- Choudhary, P. K. (2007b). A tolerance interval approach for assessment of agreement with left censored data. *Journal of Biopharmaceutical Statistics* 17(4):583–594.

- Choudhary, P. K. (2007c). Semiparametric regression for assessing agreement using tolerance bands. *Computational Statistics and Data Analysis* DOI: 10.1016/j.csda.2007.01.006
- Choudhary, P. K., Nagaraja, H. N. (2004). Measuring agreement in method comparison studies – a review. In: Balakrishnan, N., Kannan, N., Nagaraja, H. N., eds. *Advances in Ranking and Selection, Multiple Comparisons, and Reliability*. Boston: Birkhauser, pp. 215–244.
- Choudhary, P. K., Nagaraja, H. N. (2005a). Assessment of agreement using intersection-union principle. *Biometrical Journal* 47:674–681.
- Choudhary, P. K., Nagaraja, H. N. (2005b). Selecting the instrument closest to a gold standard. *Journal of Statistical Planning and Inference* 129:229–237.
- Choudhary, P. K., Nagaraja, H. N. (2005c). A two-stage procedure for selection and assessment of agreement of the best instrument with a gold-standard. *Sequential Analysis* 24:237–257.
- Choudhary, P. K., Ng, H. K. T. (2006). Assessment of agreement under non-standard conditions using regression models for mean and variance. *Biometrics* 62:288–296.
- Choudhary, P. K., Nagaraja, H. N. (2007). Tests for assessment of agreement using probability criteria. *Journal of Statistical Planning and Inference* 137:279–290.
- Cohen, J. (1960). A coefficient of agreement for nominal scales. *Educational and Psychological Measurement* 20:37–46.
- Cohen, J. (1968). Weighted kappa: Nominal scale agreement with provision for scaled disagreement or partial credit. *Psychological Bulletin* 70:213–220.
- Cronbach, L. J., Gleser, G. C., Nanda, H., Rajaratnam, N. (1972). *The Dependability of Behavioral Measurements: Theory of Generalizability for Scores and Profiles*. New York: Wiley.
- Donner, A. (1998). Sample size requirements for the comparison of two or more coefficients of inter-observer agreement. *Statistics in Medicine* 17:1157–1168.
- Dunn, D. (1989). *Statistical Evaluation of Measurement Errors: Design and Analysis of Reliability Studies*. New York: Oxford University Press Inc.
- Dunn, D. (1992). Design and analysis of reliability studies. *Statistical Methods in Medical Research* 1:123–157.
- Dunn, D. (2004). *Statistical Evaluation of Measurement Errors: Design and Analysis of Reliability Studies*. New York: Oxford University Press Inc.
- Dunn, D. (2007). Regression models for method comparison data. *Journal of Biopharmaceutical Statistics* 17(4):739–756.
- Dunn, G., Roberts, C. (1999). Modelling method comparison data. *Statistical Methods in Medical Research* 8:161–179.
- Eliasziw, M., Young, S. L., Woodbury, M. G., Fryday-Field, K. (1994). Statistical methodology for the concurrent assessment of interrater and intrarater reliability: using goniometric measurements as an example. *Physical Therapy* 74:777–788.
- Fagot, R. F. (1993). A generalized family of coefficients of relational agreement for numerical scales. *Psychometrika* 58:357–370.
- Fay, M. P. (2005). Random marginal agreement coefficients: rethinking the adjustment for chance when measuring agreement. *Biostatistics* 6:171–180.
- Fisher, R. A. (1925). *Statistical Methods for Research Workers*. Edinburgh: Oliver and Boyd.
- Fleiss, J. L. (1986). *The Design and Analysis of Clinical Experiments*. New York: Wiley.
- Food and Drug Administration (FDA) (2001). *Guidance for Industry: Bioanalytical Method Validation*. <http://www.fda.gov/cder/guidance/index.htm>
- Galton, F. (1886). Family likeness in stature. *Proceedings of the Royal Society* 40:42–73.
- Goodwin, L. D. (1997). Changing concepts of measurement validity. *Journal of Nursing Education* 36:102–107.
- Guo, Y., Manatunga, A. K. (2007). Nonparametric estimation of the concordance correlation coefficient under univariate censoring. *Biometrics* 83:164–172.

- Guttman, I. (1988). Statistical tolerance regions. In: Kotz, S., Johnson, N. L., eds. *Encyclopedia of Statistical Sciences*. Vol. 9. New York: Wiley, pp. 272–287.
- Haber, M., Barnhart, H. X. (2006). Coefficients of agreement for fixed observers. *Statistical Methods in Medical Research* 15:1–17.
- Haber, M., Barnhart, H. X. (2007). A general approach to evaluating agreement between two observers or methods of measurement. *Statistical Methods in Medical Research*, in press.
- Haber, M., Barnhart, H. X., Song, J., Gruden, J. (2005). Observer variability: a new approach in evaluating interobserver agreement. *Journal of Data Science* 3:69–83.
- Haber, M., Gao, J., Barnhart, H. X. (2007). Assessing observer agreement in studies involving replicated binary observations. *Journal of Biopharmaceutical Statistics* 17(4):757–766.
- Hand, D. J. (2004). *Measurement Theory and Practice*. New York: Oxford University Press Inc.
- Harris, I. R., Burch, B. D., St. Laurent, R. T. (2001). A blended estimator for measure of agreement with a gold standard. *Journal of Agricultural, Biological, and Environmental Statistics* 6:326–339.
- Hawkins, D. M. (2002). Diagnostics for conformity of paired quantitative measurements. *Statistics in Medicine* 21:1913–1935.
- Hutson, A. D., Wilson, D. C., Geiser, E. A. (1998). Measuring relative agreement: echocardiographer versus computer. *Journal of Agricultural, Biological and Environmental Statistics* 3:163–174.
- International Organization for Standardization (ISO) (1994). *Accuracy (Trueness and Precision) of Measurement Methods and Results—Part 1: General Principles and Definitions (5725-1)*. Geneva, Switzerland: ISO.
- Janson, H., Olsson U. (2001). A measure of agreement for interval or nominal multivariate observations. *Educational and Psychological Measurement* 61:277–289.
- Janson, H., Olsson U. (2004). A measure of agreement for interval or nominal multivariate observations by different sets of judges. *Educational and Psychological Measurement* 64:62–70.
- Kelly, G. E. (1985). Use of the structural equations model in assessing the reliability of a new measurement technique. *Applied Statistics* 34:258–263.
- King, T. S., Chinchilli, V. M. (2001a). A generalized concordance correlation coefficient for continuous and categorical data. *Statistics in Medicine* 20:2131–2147.
- King, T. S., Chinchilli, V. M. (2001b). Robust estimators of the concordance correlation coefficient. *Journal of Biopharmaceutical Statistics* 11:83–105.
- King, T. S., Chinchilli, V. M., Carrasco, J. (2007a). A repeated measures concordance correlation coefficient. *Statistics in Medicine* DOI: 10.1002/sim.2778
- King, T. S., Chinchilli, V. M., Carrasco, J. L., Wang, K. (2007b). A class of repeated measures concordance correlation coefficients. *Journal of Biopharmaceutical Statistics* 17(4):653–672.
- Krippendorff, K. (1970). Bivariate agreement coefficients for reliability of data. In: Borgatta, B. F., Bohrnstedt, G. E., eds. *Social Methodology*. San Francisco: Jossey-Bass, pp. 139–150.
- Konishi, S., Khatri, C. G., Rao, C. R. (1991). Inferences on multivariate measures of interclass and intraclass correlations in familial data. *Journal of the Royal Statistical Society Series B* 53:649–659.
- Kraemer, H. C., Periyakoil V. S., Noda, A. (2002). Kappa coefficients in medical research. *Statistics in Medicine* 21:2109–2129.
- Lee, J., Koh, D., Ong, C. N. (1989). Statistical evaluation of agreement between two methods for measuring a quantitative variable. *Computers in Biology and Medicine* 19:61–70.

- Liao, J. J. Z. (2003). An improved concordance correlation coefficient. *Pharmaceutical Statistics* 2:253–261.
- Liao, J. J. Z. (2005). Agreement for curved data. *Journal of Biopharmaceutical Statistics* 15:195–203.
- Lin, L. I. (1989). A concordance correlation coefficient to evaluate reproducibility. *Biometrics* 45:225–268.
- Lin, L. I. (1992). Assay validation using the concordance correlation coefficient. *Biometrics* 48:599–604.
- Lin, L. I. (2000a). A note on the concordance correlation coefficient. *Biometrics*, 56:324–325.
- Lin, L. I. (2000b). Total deviation index for measuring individual agreement with applications in laboratory performance and bioequivalence. *Statistics in Medicine* 19:255–270.
- Lin, L. I. (2003). Measuring Agreement. In: Chow, S., ed. *Encyclopedia of Biopharmaceutical Statistics*. Philadelphia: Informa Healthcare, pp. 561–567.
- Lin, L. I., Hedayat, A. S., Sinha, B., Yang, M. (2002). Statistical methods in assessing agreement: models, issues and tools. *Journal of American Statistical Association* 97:257–270.
- Lin, L. I., Hedayat, A. S., Wenting, W. (2007). A unified approach for assessing agreement for continuous and categorical data. *Journal of Biopharmaceutical Statistics* 17(4):629–652.
- Lin, S. C., Whipple, D. M., Ho, C. S. (1998). Evaluation of statistical equivalence using limits of agreement and associated sample size calculation. *Communications in Statistics—Theory and Methods* 27:1419–1432.
- Liu X., Du Y., Teresi J., Hasin, D. S. (2005). Concordance correlation in the measurements of time to event. *Statistics in Medicine* 24:1409–1420.
- Lord, F. M., Novick, M. R. (1968). *Statistical Theories of Mental Test Scores*. Reading, MA: Addison-Wesley.
- McGraw, K. O., Wong, S. P. (1996). Forming inferences about some intraclass correlation coefficients. *Psychological Methods* 1:30–46.
- Molenberghs, G., Vangeneugden, T., Laenen A. (2007). Estimating reliability and generalizability from hierarchical biomedical data. *Journal of Biopharmaceutical Statistics* 17(4):595–627.
- Müller, R., Büttner, P. (1994). A critical discussion of intraclass correlation coefficients. *Statistics in Medicine* 13:2465–2476.
- Nickerson, C. A. (1997) Comment on a concordance correlation to evaluate reproducibility. *Biometrics* 53:1503–1507.
- Pearson, K. (1896). VII. Mathematical contributions to the theory of evolution—III. Regression, heredity and panmixia. *Philosophical Transactions of the Royal Society, Series A* 187:253–318.
- Pearson, K. (1901). VIII. Mathematical contributions to the theory of evolution—IX. On the principle of homotypy and its relation to heredity, to the variability of the individual, and to that of the race. Part I. Homotypy in the vegetable kingdom. *Philosophical Transactions of the Royal Society, Series A* 197:285–379.
- Pearson, K., Lee, A., Bramley-Moore, L. (1899). VI. Mathematical contributions to the theory of evolution—VI. Genetic (reproductive) selection: Inheritance of fertility in man, and of fecundity in thoroughbred racehorses. *Philosophical Transactions of the Royal Society, Series A* 192:257–330.
- Raghavachari, M. (2004). Measures of concordance for assessing agreement in ratings and rank order data. In: Balakrishnan, N., Kannan, N., Nagaraja, H. N., eds. *Advances in Ranking and Selection, Multiple Comparisons and Reliability*. Boston: Birkhauser, pp. 245–263.

- Robieson, W. (1999). On Weighted Kappa and Concordance Correlation Coefficient. Ph.D. thesis, University of Illinois in Chicago.
- Quan, H., Shih, W. J. (1996). Assessing reproducibility by the within-subject coefficient of variation with random effects models. *Biometrics* 52:1195–1203.
- Quiroz, J. (2005). Assessment of equivalence using a concordance correlation coefficient in a repeated measurement design. *Journal of Biopharmaceutical Statistics* 15:913–928.
- Ryan, T.P., Woodall, W.H. (2005). The most-cited statistical papers. *Journal of Applied Statistics* 32:461–474.
- Shao, J., Zhong, B. (2004). Assessing the agreement between two quantitative assays with repeated measurements. *Journal of Biopharmaceutical Statistics* 14:201–212.
- Shavelson, R. J., Webb, N. M. (1981). Generalizability: 1973–1980. *British Journal of Mathematical and Statistical Psychology* 34:133–166.
- Shavelson, R. J., Webb, N. M. (1991). *Generalizability Theory: A Primer*. Newbury Park, CA: SAGE Publication.
- Shavelson, R. J., Webb, N. M. (1992). Generalizability theory. In: Alkin, M. C., ed. *Encyclopedia of Education Research*. Vol. 2. New York: Macmillan, pp. 538–543.
- Shavelson, R. J., Webb, N. M., Rowley, G. I. (1989). Generalizability theory. *American Psychologist* 44:922–932.
- Shoukri, M. M. (1998). Measurement of agreement. In: Armitage, P., Colton, T., eds. *The Encyclopedia of Biostatistics*. New York: Wiley, pp. 102–116.
- Shoukri, M. M., Asyali, M. H., Donner, A. (2004). Sample size requirements for the design of reliability study: review and new results. *Statistical Methods in Medical Research* 13:251–271.
- Shrout, P. E. (1998). Measurement reliability and agreement in psychiatry. *Statistical Methods in Medical Research* 7:301–317.
- Shrout, P. E., Fleiss, J. L. (1979). Intraclass correlations: uses in assessing rater reliability. *Psychological Bulletin* 86:420–428.
- St. Laurent, R. T. (1998). Evaluating agreement with a gold standard in method comparison studies. *Biometrics* 54:537–545.
- Stine, W. W. (1989). Interobserver relational agreement. *Psychological Bulletin* 106:341–347.
- Vangeneugden, T., Laenen, A., Geys, H., Renard, D., Molenberghs, G. (2004). Applying linear mixed models to estimate reliability in clinical trials with repeated measurements. *Controlled Clinical Trials* 25:13–30.
- Vangeneugden, T., Laenen, A., Geys, H., Renard, D., Molenberghs, G. (2005). Applying concepts of generalizability theory on clinical trial data to investigate sources of variation and their impact on reliability. *Biometrics* 61:295–304.
- Wang, W., Hwang, J. R. G. (2001). A nearly unbiased test for individual bioequivalence problems using probability criteria. *Journal of Statistical Planning and Inference* 99:41–58.
- Williamson, J. M., Crawford, S. B., Lin, H. (2007). Resampling dependent concordance correlation coefficients. *Journal of Biopharmaceutical Statistics* 17(4):685–696.
- Zhong, B., Shao, J. (2003). Evaluating the agreement of two quantitative assays with repeated measurements. *Journal of Biopharmaceutical Statistics* 13:75–86.
- Zegers, F. E. (1986). A family of chance-corrected association coefficients for metric scales. *Psychometrika* 51:559–562.