Interval Methods for Data Fitting under Uncertainty: A Probabilistic Treatment

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Abstract

How to estimate parameters from observations subject to errors and uncertainty? Very often, the measurement errors are random quantities that can be adequately described by the probability theory. When we know that the measurement errors are normally distributed with zero mean, then the (asymptotically optimal) Maximum Likelihood Method leads to the popular least squares estimates. In many situations, however, we do not know the shape of the error distribution, we only know that the measurement errors are located on a certain interval. Then the maximum entropy approach leads to a uniform distribution on this interval, and the Maximum Likelihood Method results in the so-called minimax estimates. We analyze specificity and drawbacks of the minimax estimation under essential interval uncertainty in data and discuss possible ways to solve the difficulties. Finally, we show that, for the linear functional dependency, the minimax estimates motivated by the Maximum Likelihood Method coincide with those produced by the Maximum Compatibility Method that originate from interval analysis.

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1 Introduction

The paper is devoted to data analysis under uncertainty, when we do not know exact values of our measurements, observations, etc., but instead we have at our disposal some alternative information. These can be bounds of possible errors of the measured quantities, both lower and upper, which is equivalent to determining interval results of measurements. On the other hand, these can be probabilistic characteristics of the errors corrupting the measured quantities, and then we can use well-developed methods of probability theory to process our data.

Data analysis methods based on probability theory are very elaborate and popular, they have been applied to processing the results of measurements and observations for about two centuries. Interval methods came into being in the middle of the XX century, and nowadays they only start winning the favor of engineers and practitioners. The purpose of our paper is to show that the results and conclusions drawn by these different approaches are in good conformity with each other, so that any one of them can be used to justify and substantiate (or even to verify) the other.

Interval methods in data analysis originated from the pioneering works by L. V. Kantorovich[9] and F. C. Schweppe [24]. For the last decades, they have been developed deeply and extensively by many researchers throughout the world.

Broad research area and a great variety of applications resulted in great terminological diversity: doing similar (and even the same) things, people speak of “guaranteed parameter estimation”, “set-membership estimation”, “bounded-error approach”, “interval data fitting”, “interval regression”, etc. To get an insight into the current state of knowledge in this field, the reader can consult, e.g., [7, 13] and the literature cited there.

Our paper appears in the special issue of the journal “Reliable Computing” devoted to Ramon E. Moore and his scientific heritage. The topic of the paper has a direct relationship to the activity of Ramon Moore who suggested, in [15], an elegant way to present and describe the solution sets in nonlinear parameter estimation.

Formulation of the problem. In many situations, we know the general form of the functional dependency between the quantities \( x = (x_1, \ldots, x_k) \) and \( y \), i.e., we know that

\[
y = f(x, p),
\]

where \( p = (p_1, \ldots, p_l) \) is an \( l \)-dimensional parameter vector. Based on actual values of the variables \( x \) and \( y \), we have to find the values of \( p_1, \ldots, p_l \) that correspond to a specific function \( f \) from the parametric family [1]. This problem is referred to as “parameter estimation problem”, “data fitting problem”, “regression problem”, “curve fitting”, etc.

\[1\]Leonid Kantorovich was one of the founders, along with George Dantzig, of the linear programming, as well as Nobel Prize winner in economic sciences.
To find the parameter $p$, we repeatedly measure the corresponding values of $x$ and $y$. As a result, for each measurement $i = 1, 2, \ldots, m$, we get the corresponding values $x^{(i)} = (x_{1}^{(i)}, \ldots, x_{k}^{(i)})$ and $y^{(i)}$. Our task is to determine such $p^* = (p_1^*, \ldots, p_l^*)$ that the function $y = f(x, p^*)$ “best fits” the measurement data set

\[
\begin{align*}
  &x^{(1)}, \; y^{(1)}, \\
  &x^{(2)}, \; y^{(2)}, \\
  &\cdots, \; \cdots, \\
  &x^{(m)}, \; y^{(m)}
\end{align*}
\]  

(see Fig. 1). At this point, we have to explain the sense in which we understand the “best fitting”.

![Figure 1: In data fitting problem, we have to construct a line that best fits measurement data.](image)

Ideally, the “best fit” line should go through all the measurement points of the set $\{2\}$. This happens when the measurements are so accurate that we can safely ignore any errors and assume that the measured values of the quantities $x_1, x_2, \ldots, x_k$, and $y$ are exact. In this case, we get a system of equations

\[
\begin{align*}
  f(x^{(1)}, p) &= y^{(1)}, \\
  f(x^{(2)}, p) &= y^{(2)}, \\
  \vdots &= \vdots, \\
  f(x^{(m)}, p) &= y^{(m)},
\end{align*}
\]  

(3)

with $l$ unknowns $p_1, \ldots, p_l$. Having solved the system, we obtain the desired values of the parameters that correspond to the data analyzed. In general, to find $l$ unknowns, it is sufficient to have $l$ different equations. Thus, in the ideal case of absolutely accurate measurements and providing that $f$ adequately
describes the actual functional dependency under study, we would be able to
determine the values of all \( l \) parameters \( p_i \) after \( m = l \) measurements.

However, measurements are not precise. They can even include so-called
outliers, measurement results that largely deviate from the rest of the data
set, which may be caused by experimental errors, etc. The function \( f \) may
not perfectly reflect the functional dependency that exists between \( x \) and \( y \) in
practice. Usually, \( f \) is considered as a “main part” (to within neglected terms)
of the actual function between \( x \) and \( y \). And so on. In such situations, we
cannot expect that the equalities \( y^{(i)} = f(x^{(i)}, p), i = 1, 2, \ldots, m, \) are exactly
satisfied for some \( p \). Then the “best fit” line should be a best approximation, in
a prescribed sense, of the data set (2) (this is the situation described at Fig. 1).

In such situations, the equation system (3) cannot be solved exactly, and
we have to find its solution in some generalized sense. The latter is especially
truer since we often try to make as much measurements as possible: every mea-
surement provides us with additional information about \( f \). Hence, the equation
system (3) can be over-determined.

We are going to consider a simplified situation with the measurement errors:
the values \( x^{(i)} \) of the argument are supposed to be exact, but the actual values
\( y^{(i)} \) of the function \( f \), i.e., such that \( y^{(i)} = f(x^{(i)}, p) \), are, in general, different
from the measured values of \( f \), denoted by \( \tilde{y}^{(i)} \). So, instead of (2), we have to
process the approximate data

\[
\begin{align*}
  x^{(1)}, & \quad \tilde{y}^{(1)}, \\
  x^{(2)}, & \quad \tilde{y}^{(2)}, \\
  \ldots, & \quad \ldots, \\
  x^{(m)}, & \quad \tilde{y}^{(m)},
\end{align*}
\]

which results in the necessity to solve the equation system

\[
\begin{align*}
  f(x^{(1)}, p) &= \tilde{y}^{(1)}, \\
  f(x^{(2)}, p) &= \tilde{y}^{(2)}, \\
  \vdots & \quad \vdots \\
  f(x^{(m)}, p) &= \tilde{y}^{(m)},
\end{align*}
\]

rather than system (3).

\[ \text{2 Probabilistic approaches} \]

In this section, we discuss how to estimate parameters when our measurements
are not exact, but we have certain probabilistic information about measurement
errors. The material we present below is quite standard, but we survey it
to make our paper self-sufficient, as well as to prepare and motivate further
conclusions.
Notice that even the fact that probabilistic approaches are applicable to processing specific data is not trivial and should be substantiated on its own. In particular, the so-called statistical stability (see [5]) is one of necessary prerequisites for the methods of probability theory in statistics to be adequate and efficient.

**Typical probabilistic prerequisites.** In some cases, we can be sure that the probability theory is an adequate tool for describing the measurement errors. Moreover, the probability distributions of the measurement errors \( \Delta y^{(i)} = \tilde{y}^{(i)} - y^{(i)} \) or at least the shape of the corresponding probability distributions are known. Let us consider this kind of situation.

First of all, from the repeated measurements, we often know, with good accuracy, the mean value of the measurement error. In such situations, we can correct the measurement results by subtracting this bias (mean value). So, we can suppose that the mean value of the measurement error is equal to 0.

Frequently, when all major measurement disturbances are eliminated, the remaining errors are formed from many small error sources. Under reasonable conditions, the probability distribution of the sum of \( N \) small independent random variables is known to tend to the normal distribution when \( N \) increases. This fact is called the Central Limit Theorem (see, e.g., [22, 33]). Thus, when the measurement error comes from the joint effect of a large number of small independent components, we can safely assume that the resulting probability distribution of the overall measurement error is normal. For a general normal Gaussian distribution with zero mean, the probability density \( \rho \) as a function of the error \( \Delta y \) has the form

\[
\rho(\Delta y) = \frac{1}{\sigma \sqrt{2\pi}} \cdot \exp\left(\frac{-(\Delta y)^2}{2\sigma^2}\right)
\]  

for an appropriate value \( \sigma \) (called standard deviation).

We would remind the following empirical fact: in practice, about 60% of the measuring instruments have a normally distributed measurement error; see, e.g., [19, 20].

**How to estimate the parameters in the case of normal error distribution.** It is well known that an asymptotically optimal way to determine the parameters of a distribution from the sample values is the Maximum Likelihood Method. According to it, we select the values of the unknown parameters so that the probability of the given sample of values (in the case of discrete probability distributions) or probability density at the given sample of values (in the case of continuous probability distributions) are the largest possible; see, e.g., [11, 14].

Measurement errors corresponding to different measurements are usually assumed to be independent. The probability of several independent events occurring together is equal to the product of the corresponding probabilities. Thus, for the normal distribution (6), the above implies that we have to select the
unknown parameters $p$ that maximize the product

$$L = \prod_{i=1}^{m} \rho(\Delta y^{(i)}) = \prod_{i=1}^{m} \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(\Delta y^{(i)})^2}{2\sigma^2}\right).$$

Substituting the expression $\Delta y^{(i)} = \tilde{y}^{(i)} - f(x^{(i)}, p)$ into this formula, we conclude that $p$ should maximize the product

$$L = \prod_{i=1}^{m} \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(	ilde{y}^{(i)} - f(x^{(i)}, p))^2}{2\sigma^2}\right).$$

The constant coefficient $1/(\sigma\sqrt{2\pi})$ is the same for all $i$, so we have

$$L = \frac{1}{(\sigma\sqrt{2\pi})^m} \prod_{i=1}^{m} \exp\left(-\frac{(	ilde{y}^{(i)} - f(x^{(i)}, p))^2}{2\sigma^2}\right)$$

$$= \frac{1}{(\sigma\sqrt{2\pi})^m} \exp\left(-\frac{1}{2\sigma^2} \sum_{i=1}^{m} \left(\tilde{y}^{(i)} - f(x^{(i)}, p)\right)^2\right).$$

From monotonicity of the function $\exp$, it follows that maximizing $L$ is equivalent to minimizing the sum

$$\sum_{i=1}^{m} \left(\tilde{y}^{(i)} - f(x^{(i)}, p)\right)^2. \tag{7}$$

We thus arrive at the well-known least squares method independently proposed by A.-M. Legendre and K. F. Gauss (see, e.g., [6, 11]): the value of the parameter $p$ under estimation should be selected so that the sum of the squares of the approximation errors $\Delta y^{(i)} = \tilde{y}^{(i)} - f(x^{(i)}, p)$ is the smallest possible. Using the least squares method is an asymptotically optimal way to estimate the parameters from the observations when the measurement errors are normally distributed with zero mean [6, 11].

Need to consider situations where the error distribution is not known, in particular, situation of interval uncertainty. The above approach and the least squares method are not universal. As we have mentioned, for about 40% of measuring instruments, the measurement errors are not normally distributed. Moreover, often, we do not have enough measurements to determine the actual shape of the corresponding probability distribution, which is known as “small sample size” problem.

In many such situations, the only information we know is that the possible values of the measurement error are located within some bounds $\Delta$ and $\tilde{\Delta}$, but we do not know the probability of different values within the interval $[\Delta, \tilde{\Delta}]$; see numerous examples in [22].

Such is the situation with many real-life sensors. In principle, we can compare the results of using a given sensor with the results of measuring the same
quantity by a super-accurate measuring instrument and thus, get the desired probability distribution of the measurement error. However, such a calibration is usually very expensive — often several orders of magnitude more expensive that the price of the sensor itself. In such cases, the only information that we have about the measurement error is the upper bound on its absolute value — this information is usually provided by the manufacturer of the corresponding measuring instrument.

Instead of calibrating each individual measuring instrument, practitioners often simply compose it of several components with known characteristics and known tolerances, i.e., known bounds on the difference between the actual and the nominal values of these characteristics. In this case, we only know the bounds on the corresponding values of the measured quantity, that is, only bounds on the measurement error.

What shall we do in this case?

How to estimate the parameters when the error distribution is not known. Situations when we do not know the exact probabilistic distribution of the measurement errors are ubiquitous. Then, several different distributions can be consistent with our knowledge. Some of these distributions are more informative, some are less. It is reasonable to select, among all the distributions, the one with the least possible amount of information, i.e., the one which does not add anything to our knowledge that the random variable is located on a given interval.

If a random variable has the probability density function \( \rho(x) \), the amount of information it bears is usually described by the entropy

\[
S = - \int_{-\infty}^{\infty} \rho(x) \ln \rho(x) \, dx
\]  

(see, e.g., \[2, 8, 17\]). In fact, the entropy gives a measure of how chaotic the probabilistic distribution is. So, the more entropy of a distribution, the less informative it is. Then the above idea means that, among all the probability distributions \( \rho(x) \) consistent with our knowledge, we should select the one for which entropy (8) attains the largest possible value. This consideration is known as the maximum entropy approach.

Let us apply the maximum entropy approach to the case when the only information about the probability distribution is that it is located on the interval \([\Delta, \bar{\Delta}]\), i.e., that \( \rho(x) = 0 \) for \( x \notin [\Delta, \bar{\Delta}] \). Thus, we need to maximize the entropy

\[
S = - \int_{-\infty}^{+\infty} \rho(x) \ln \rho(x) \, dx = - \int_{\Delta}^{\bar{\Delta}} \rho(x) \ln \rho(x) \, dx.
\]  

(9)

Additionally, expression (9) should be maximized under the constraint that the overall probability is 1, i.e.,

\[
\int_{\Delta}^{\bar{\Delta}} \rho(x) \, dx = 1.
\]  

(10)
The method of Lagrange multipliers reduces this constraint optimization problem to an unconstrained problem of maximizing the function
\[- \int_\Delta^\Sigma \rho(x) \ln \rho(x) \, dx + \lambda \left( \int_\Delta^\Sigma \rho(x) \, dx - 1 \right),\]
where \(\lambda\) is a Lagrange multiplier. Taking the functional (variational) derivative over \(\rho(x)\) (see [4]) and equating this derivative to 0, we conclude that
\[- \ln \rho(x) - 1 + \lambda = 0,\]
hence \(\ln \rho(x) = \lambda - 1 = \text{const}\) and thus, \(\rho(x) = \text{const}\). The value of this constant can be determined from equality [10] that the overall probability should be equal to 1. Therefore, we have a uniform distribution with the probability density
\[\rho(x) = \frac{1}{\Sigma - \Delta} = \text{const}. \tag{11}\]

The conclusion that we should select a uniform distribution is in good accordance with common sense. Indeed, since we have no reason to believe that some values from the interval \([\Delta, \Sigma]\) are more probable and some values are less probable, it makes sense to select a distribution in which all these values are equally probable, i.e., a uniform distribution on this interval.

As we have mentioned, it is usually reasonable to assume that the main part of the systematic error (bias) has already been eliminated and thus, the mean value for the measurement error is 0. For a uniform distribution, the mean value is equal to the midpoint of the corresponding interval, and the above requirement takes the form
\[\frac{\Delta + \Sigma}{2} = 0.\]
So, if we denote \(\Delta \overset{\text{def}}{=} \overline{\Delta}\), then \(\Delta = -\Delta\), and the interval should be \([-\Delta, \Delta]\) for a certain value \(\Delta\).

**How to estimate the parameters in the case of uniform error distribution.** In this situation, we have \(l + 1\) unknowns: \(l\) parameters \(p = (p_1, \ldots, p_l)\) and the parameter \(\Delta\) that characterizes the measurement uncertainty. To find the parameters, we are going to use, similar to the case of normal error distribution, the Maximum Likelihood Method, i.e., we maximize the product
\[\mathcal{L} = \prod_{i=1}^{m} \rho(\Delta y^{(i)}).\]

The probability distribution \(\rho(x)\) is located on the interval \([-\Delta, \Delta]\). Therefore, if the absolute value \(|\Delta y^{(i)}|\) of one of the measurement errors \(\Delta y^{(i)}\) exceeds \(\Delta\),

\[\text{The mathematical fact that the uniform distribution provides the maximum entropy is widely known, being a “mathematical folklore”, and its proof can be found in many sources, e.g., in [17].}\]
then the corresponding factor $\rho(\Delta y^{(i)})$ equals 0 and, consequently, the entire product $\mathcal{L}$ is equal to 0 too.

To find the desired maximum of $\mathcal{L}$, we have to consider only the values $p$ and $\Delta$ for which

$$|\Delta y^{(i)}| \leq \Delta, \quad i = 1, 2, \ldots, m,$$

i.e., for which $\Delta \geq \max_i |\Delta y^{(i)}|$. For such values, $\rho(\Delta y^{(i)}) = 1/(2\Delta)$ according to (11), and the product $\mathcal{L}$ takes the form

$$\mathcal{L} = \frac{1}{(2\Delta)^m}.$$  

This expression is the largest if and only if the corresponding value $\Delta$ is the smallest.

For fixed $p$, the only restriction on $\Delta$ is that

$$\Delta \geq \max_{1 \leq i \leq m} |\Delta y^{(i)}| = \max_{1 \leq i \leq m} |\tilde{y}^{(i)} - f(x^{(i)}, p)|.$$  

Thus, for each tuple $p$, the smallest possible $\Delta$ is the smallest value that satisfies this inequality, i.e., it is the value

$$\Delta = \max_{1 \leq i \leq m} |\tilde{y}^{(i)} - f(x^{(i)}, p)|. \quad (13)$$

We want to select the parameters $p$ for which the probability $\mathcal{L}$ is the largest, i.e., equivalently, for which the value $\Delta$ described by formula (13) is the smallest one. Thus, in situations where the shape of the probability distribution of the measurement error is not known, we should select the parameters $p$ for which expression (13) is the smallest possible, i.e., for which

$$\min_p \max_{1 \leq i \leq m} |\tilde{y}^{(i)} - f(x^{(i)}, p)| \quad (14)$$

is attained.

The above result, expressed by formulas (13)–(14), is not quite trivial, and it makes sense to comment on it. In principle, estimating the parameters $p = (p_1, p_2, \ldots, p_l)$ of the unknown functional dependency $y = f(x, p)$ that best fit the given data amounts to computing, for the equation system (5), a “pseudo-solution” that minimizes the discrepancy between the left-hand and right-hand sides of (5). To do that, we construct the so-called defect vector $d \in \mathbb{R}^m$ which is made up of the differences $d_i = (\tilde{y}^{(i)} - f(x^{(i)}, p))$, $i = 1, 2, \ldots, m$, between the left-hand and right-hand sides of system (5), and then make it as small as possible. In fact, this means that we take a norm of the defect vector, as a general measure of how large it is, and then minimize this norm over $p$. If the minimum of the norm of the defect equals zero, we get an exact solution to system (5). Otherwise, a pseudo-solution is obtained that corresponds to approximation of the data set by the best-fit line (the situation depicted at Fig. 1).
A norm of the defect vector can be constructed from the defects of separate components \( d_i = (\tilde{y}(i) - f(x(i), p)), \) \( i = 1, 2, \ldots, m, \) in various ways. In particular, taking the Euclidean norm (2-norm) of the defect, i.e.,

\[
\|d\|_2 = \left( \sum_{i=1}^{m} d_i^2 \right)^{1/2},
\]

results in (7) and corresponds to the least squares method. Our reasoning above shows that only the Chebyshev norm (\( \infty \)-norm), i.e.,

\[
\|d\|_\infty = \max_{1 \leq i \leq m} d_i,
\]

has a clear probabilistic interpretation when we do not know of any information about the probability density functions of the error. Formula (13) expresses the essence of popular “minimax estimation” (see, e.g., [7, 11, 35]), and it is often used without any probabilistic context due to its clear meaning.

To conclude, it is worth noting that the experience of applying the Maximum Likelihood Method for joint considering interval and probabilistic approaches in the data fitting problems is not new. S. Zhilin in [37] used the maximum likelihood estimation in experimental study of the correlation between different types of estimates. The conclusion made in [37] on the base of extensive numerical simulation is that the non-probabilistic interval-based estimates are very good and even the best ones for uniform or nearly uniform probabilistic distributions.

3 Specific features of interval approaches

In interval approaches to the data fitting problem, we assume that the only information we have about each measurements error \( \Delta y(i) \) is the upper bound on its absolute value: \( |\Delta y(i)| \leq \Delta. \) In this case, the only information that we have about each actual (unknown) value \( y(i) = f(x(i), p) \) is that this value belongs to the interval

\[
y(i) \overset{\text{def}}{=} [\tilde{y}(i) - \Delta, \tilde{y}(i) + \Delta], \quad i = 1, 2, \ldots, m,
\]

Then, the inequalities (12)

\[
|\Delta y(i)| = |\tilde{y}(i) - f(x(i), p)| \leq \Delta
\]

can be equivalently rewritten as

\[
f(x(i), p) \in y(i) \quad i = 1, 2, \ldots, m.
\]

In this case, the sense of the parameter estimation is depicted at Fig. 2: we have to construct a line from a given parametric family that “best fits” the interval data in a prescribed sense. Notice that such “best fit line” may go through each segment representing data uncertainty or, alternatively, may not do that. In the
former case, when the constructed line intersects all the uncertainty intervals \((x_i, y^{(i)})\), \(i = 1, 2, \ldots, m\), we will speak that the parameters of the line are compatible with the interval data.

Overall, instead of the equation system (5), we arrive at an analogous system of interval equations

\[
\begin{align*}
  f(x^{(1)}, p) &= y^{(1)}, \\
  f(x^{(2)}, p) &= y^{(2)}, \\
  &\vdots \ddots \vdots \\
  f(x^{(m)}, p) &= y^{(m)},
\end{align*}
\]

that we have to “solve” with respect to \(p\) for parameter estimation.

![Figure 2: Data fitting problem for interval measurement data: we have to construct a line that best fits the data.](image)

In modern interval analysis, there exist several concepts of “solution” and “solution sets” to interval systems of equations, and the most relevant to our task is the so-called “united solution set”. It is defined as the set of all solutions to the usual point systems of the same form with the parameters taken from the prescribed intervals. For (16), the united solution set is formally determined as

\[
\Xi = \{ p \in \mathbb{R}^n \mid \exists y^{(1)} \in y^{(1)} \cdots \exists y^{(m)} \in y^{(m)} \}
\]

being made up of solutions to the usual (point) systems of the form (3) with \(y^{(1)}, y^{(2)}, \ldots, y^{(m)}\) from the intervals \(y^{(1)}, y^{(2)}, \ldots, y^{(m)}\). If the solution set \(\Xi\) is not empty, it describes the set of parameters compatible with the measurement.
and each \( p \in \Xi \) can be taken as a solution to the data fitting problem, i.e., as an estimate of the parameters. Still, set (17) may not coincide with the set of solutions to the data fitting problem in general, owing to additional requirements on the parameter estimates that can be present in the problem statement. This is why we are going to use, with respect to (17), the term feasible parameter set, popular in the set-membership estimation theory.

The Demidenko paradox. The case of nonempty feasible parameter set \( \Xi \) is the most favorable in interval data fitting, but we do not have to reject the opposite case when the set \( \Xi \) is empty. Then there does not exist \( p \) strictly compatible with the data set (18), but this in no way means that the data fitting problem is not solvable. This only means that its solution has the other status. For better understanding our reasoning, it is worth reminding that for usual non-interval data fitting problem, which is the limit case of the interval problem statement, the compatibility between the data and parameters in the sense of our definition is mostly unachieved, being a very rare and exceptional event.

Moreover, if possible emptiness of the feasible parameter set is not taken into account, we can come to a paradox that was first noticed by E. Demidenko in the note [3]. Its essence can be described in short by the phrase “the better, the worse”. Specifically, the more accurate are our measurements and more narrow the uncertainty intervals \( y^{(1)}, y^{(2)}, \ldots, y^{(m)} \), the more likely that the feasible parameter set \( \Xi \) is empty and the data fitting problem looks “unsolvable”. Conversely, if we organize crude measurement and our errors are large, the intervals \( y^{(1)}, y^{(2)}, \ldots, y^{(m)} \) are wide, but this enlarges the feasible parameter set \( \Xi \), and it is easier to take an estimate of the parameters from it.

In fact, the Demidenko paradox is based on the assumptions that

(i) the intervals \( y^{(1)}, y^{(2)}, \ldots, y^{(m)} \) represent rigorous bounds on the actual values of our measurements and falling outside them is impossible,

(ii) the function \( f \) exactly represents the functional dependency between \( x \) and \( y \).

Any one or both of the above points can be violated in real life problems. For example, we can consider the intervals \( y^{(1)}, y^{(2)}, \ldots, y^{(m)} \) as “soft” bounds on the respective measurements: these bounds are not obligatory for the actual values of \( y^{(1)}, y^{(2)}, \ldots, y^{(m)} \), but rather serve as estimates of their ranges. Our data can have outliers that spoil the ideal measurements picture. The function \( f \) may be an approximation to the actual dependency between \( x \) and \( y \) taken
Bad accuracy and wide uncertainty intervals enable one to construct many models compatible with the data:

For better accuracy and narrow uncertainty intervals, there might not exist a model compatible with the data:

Figure 3: The essence of the Demidenko paradox.

(e.g., from physical, chemical, economic, etc., reasons) merely as a dominant trend. And so on.

Therefore, the emptiness of the feasible parameter set should not be perceived as a signal for terminating the solution of the data fitting problem. We have to accept the possibility for the feasible parameter set to be empty, the possibility that there are no parameters strictly compatible with the processed data. Even more, our approach to the data fitting and parameter estimation for interval data should successfully cope with this “double-layer character” of the problem, providing a uniform treatment of the two situations when the feasible
parameter set is either empty or nonempty. This is a specific feature of the data processing under essentially interval uncertainty.

**Minimax estimation for interval data.** Let us turn to the minimax estimation (13)–(14) derived from the Maximum Likelihood Method in the previous section.

Inclusions (15) mean that if the upper bound on the absolute value of the measurement error is equal to $\Delta$, then the actual value $f(x^{(i)}, p)$ of the quantity $y$ agrees with the measurement result $\tilde{y}^{(i)}$. In these terms, expression (13) is the smallest bound $\Delta$ for which all the measurement results agree with the actual values; the selected value $p$ is the one for which the difference between the measurement results and the actual values is the smallest possible. Overall, a natural probabilistic approach to estimating parameters, the Maximum Likelihood Method, leads to an “interval-ready” formula (14) for parameter estimation, which can be directly applied to data fitting under interval uncertainty. However, some points should be tuned to this specific case.

Under certain conditions, we may have a priori bounds on some parameters $p_j$, i.e., the values $\underline{p}_j$ and $\overline{p}_j$ for which the actual (unknown) value $p_j$ satisfies the inequality $\underline{p}_j \leq p_j \leq \overline{p}_j$. Then we should minimize expression (13) for $\Delta$ only over the tuples $p = (p_1, \ldots, p_l)$ which satisfy these inequalities.

Very often, we can be interested only in some of the parameters. Without losing generality, let us assume that we have to find the values of the parameters $p_1, \ldots, p_\ell$ for some $\ell < l$, while the parameters $p_{\ell+1}, \ldots, p_l$ are not interesting for us. Then we can use the mathematical fact that

$$
\min_{p_1, p_2, \ldots, p_{\ell}} \max_{1 \leq i \leq m} \left| \tilde{y}^{(i)} - f(x^{(i)}, p) \right| = \min_{p_1, p_2, \ldots, p_\ell} \min_{p_{\ell+1}, \ldots, p_l} \max_{1 \leq i \leq m} \left| \tilde{y}^{(i)} - f(x^{(i)}, p) \right|.
$$

In other words, minimizing expression (13) over all the tuples $(p_1, \ldots, p_l)$ is equivalent to minimizing an auxiliary expression

$$
\Phi(p_1, p_2, \ldots, p_\ell) \overset{\text{def}}{=} \min_{p_{\ell+1}, \ldots, p_l} \max_{1 \leq i \leq m} \left| \tilde{y}^{(i)} - f(x^{(i)}, p) \right|
$$

over the parameters $p_1, \ldots, p_\ell$. This trick may prove very helpful in situations when we want to convert some uncertain factors into parameters.

**Plateau-like extremum region.** Yet another specific feature of the data fitting problem under essential interval uncertainty is that its feasible parameter set $\Xi$ may be a “real” non-singleton set made up of infinitely many points. In fact, such feasible parameter sets can have nonempty interior and nonzero measure, so that the solution to the minimization problem (19) is achieved at a whole region in $\mathbb{R}^{\ell}$. In mathematical terms, this feature is expressed by the fact that the feasible parameter set, i.e., the united solution set to the interval equation system (16) that we have to solve for constructing the best fit line, is a
solid set. For every point of such feasible parameter set, the norm of the defect
\[ \max_i |\tilde{y}(i) - f(x(i), p)| \] is precisely zero. So, the objective function (19) attains
its minimum at a zero level plateau like that depicted at the right-hand side
of Fig. 4. That is very disadvantageous for our minimization problem, both in
itself and due to practical reasons.

First of all, checking whether a point belongs to the feasible parameter set
amounts to testing unstable equality of the objective function (19) to zero.
Furthermore, the points of the feasible parameter set \( \Xi \) are, in reality, different
from each other with respect to the data fitting problem. In particular, it makes
sense to distinguish between points of the boundary and interior of the set \( \Xi \),
since the latter are stable under data perturbations. Choosing the interior
points of the feasible parameter set \( \Xi \) as parameter estimates would be more
preferable, but the objective function (19) does not allow us to do that.

Figure 4: Various configurations of the minima
in (14) and (19) depending on their signs.

By and large, it is intuitively clear that the estimates taken somewhere
in the “middle” of the feasible parameter set \( \Xi \) are “more robust” and even
“more probable”. This is confirmed, for example, by computational experiments
from [26] demonstrating that the uniform probabilistic distributions over input
variables of a function lead to a non-uniform distribution over its range of values,
with the maximum probability density in the central area.

Figure 5: Desirable configuration of the negative minimum.

The situation with the zero-level plateau in the minimax estimation (Fig. 4,
right) should be corrected, and some parameter estimation methods really do
that during their execution. For example, in the uncertainty center method from \cite{21, 37, 38}, the estimate is taken as the middle point of the outer box for the feasible parameter set. Ideally, it would be desirable to have the configuration of the minimum for (19) like that depicted at Fig. 5, where the objective function helps distinguishing the interior points from the feasible parameter set. In Section (5), we are going to show how one can naturally improve, in the above sense, the minimax estimation for interval data fitting with the linear functional dependency.

4 Straight line fitting

As an example of applying the ideas developed in the preceding sections, let us consider the problem of solving an $m \times n$-system of interval linear equations that arises in the interval data fitting problem for linear functional dependency. It is often called “linear regression problem”, etc.

Suppose that we are given a system of linear equations

$$
\sum_{j=1}^{n} a_{ij} z_j - b_i = 0, \quad 1 \leq i \leq m,
$$

under interval uncertainty, when we only know that $a_{ij}$ and $b_i$ belong to intervals $a_{ij} = [a_{ij}, \overline{a}_{ij}]$ and $b_i = [\underline{b}_i, \overline{b}_i]$ respectively. One can think that an interval $m \times n$-matrix $A = (a_{ij})$ and an interval $m$-vector $b = (b_i)$ are given that specify an interval linear system of algebraic equations $Az - b = 0$, which is equivalent to $A z = b$. We are interested in the values of the variables $z_1, \ldots, z_n$ that “best fit” equalities (20) under the uncertainty represented by the data set $a_{ij}, b_i, i = 1, \ldots, m, j = 1, \ldots, n$ (see Fig. 6).

The problem is in close relationship with the general data fitting problem we have considered in the previous sections. One can see here $m$ observations based on which we would like to determine the desired parameters $z_1, z_2, \ldots, z_n$. As for the interval data $a_{ij}, b_i, i = 1, \ldots, m, j = 1, \ldots, n$, we will not consider them as measurement results, but represent them as additional parameters using the technique elaborated at the end of the previous section. So, in our problem statement, the number $\ell$ of the sought-for parameters is equal to $n$, and these parameters $p_1, p_2, \ldots, p_\ell$ from formulation (19) coincide with $z_1, z_2, \ldots, z_n$ from (20). In addition to these $\ell = n$ parameters of interest, we also have the following auxiliary parameters:

- $m \cdot n$ parameters $a_{ij}, 1 \leq i \leq m, 1 \leq j \leq n,$
  - for which we are given the bounds $a_{ij}$ and $\overline{a}_{ij}$;
- $m$ parameters $b_i, 1 \leq i \leq m,$
  - for which we are given the bounds $\underline{b}_i$ and $\overline{b}_i$.

In total, we consider $mn + m + n$ parameters, which can be represented as a long composite vector $p = (z, A, b), A = (a_{ij}), b = (b_i)$. As an input $x^{(i)}$ of the function that describes the $i$-th observation, we can simply take $x^{(i)} = i$. 
Overall, the model-based value $f(x, p)$ of the observed functional dependency is exactly

$$f(x^{(i)}, p) = f(i, z, A, b) = \sum_{j=1}^{n} a_{ij} z_j - b_i,$$

while the observed value $y^{(i)}$ is always equal to 0.

For this problem, formulation (19) originated from the Maximum Likelihood Method leads to selecting the values $z_1, \ldots, z_n$ for which the following expression is minimized:

$$\Phi(z) = \max_{1 \leq i \leq m} \min_{a_{ij} \in a_{ij}, b_i \in b_i} \left| \sum_{j=1}^{n} a_{ij} z_j - b_i \right|.$$  

Here, each of the parameters $a_{ij}, j = 1, 2, \ldots, n$, and $b_i$ occurs only in the $i$-th expression under $\min_{1 \leq i \leq m}$. So, minimizing $\max_{1 \leq i \leq m}$ is equivalent to minimizing the corresponding expression for each $i$:

$$\Phi(z) = \max_{1 \leq i \leq m} \min_{a_{ij} \in a_{ij}, b_i \in b_i} \left| \sum_{j=1}^{n} a_{ij} z_j - b_i \right|. \quad (21)$$

Next, the inner minimum in (21) can be found explicitly. The range of values of the linear expression $\sum_{j=1}^{n} a_{ij} z_j - b_i$ over all $a_{ij} \in a_{ij}$ and $b_i \in b_i$ coincides with its natural interval extension due to the fundamental theorem of interval arithmetic [16, 18, 31]. Therefore,

$$\min_{a_{ij} \in a_{ij}, b_i \in b_i} \left| \sum_{j=1}^{n} a_{ij} z_j - b_i \right| = \left\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \right\rangle,$$
where \( \langle \cdot \rangle \) means mignitude of the interval (see [18, 31]), the smallest distance from points of the interval to zero:

\[
\langle u \rangle = \begin{cases} 
\min\{|u|, |u|\}, & \text{if } 0 \not\in u, \\
0, & \text{otherwise}.
\end{cases}
\]

Overall, we arrive at the following optimization problem:

\[
\text{find } \min_{z_1, z_2, \ldots, z_n} \Phi(z_1, z_2, \ldots, z_n),
\]

where

\[
\Phi(z) = \Phi(z_1, z_2, \ldots, z_n) = \max_{1 \leq i \leq m} \left\{ \sum_{j=1}^{n} a_{ij} z_j - b_i \right\}.
\]

We would remind that, for interval systems of linear algebraic equations \( Az = b \), the \textit{united solution set} (also called just \textit{solution set}) is defined, similar to (17), as the set of all solutions to the point systems \( Az = b \) with \( A \in A \) and \( b \in b \). In formal terms, the united solution set to \( Az = b \) is

\[
\Xi(A, b) \overset{\text{def}}{=} \left\{ z \in \mathbb{R}^n \mid (\exists A \in A)(\exists b \in b)(Az = b) \right\}.
\]

As before, the set \( \Xi(A, b) \) is the “feasible parameter set” for our data fitting problem under interval uncertainty, i.e., the set where there is strict compatibility between the data and the parameters of the straight line.

If the set \( \Xi(A, b) \) is empty, then the minimum in the optimization problem (22)–(23) is strictly positive, which corresponds to the left-hand side picture at Fig. 4. A specific feature of our situation is that the objective function \( \Phi(z) \) is non-smooth (piecewise linear), and its minimum is attained at a sharp point of its graph. If the set \( \Xi(A, b) \) is nonempty, then the minimum in (22) is zero, which corresponds to the right-hand side picture at Fig. 4. The minimum is then attained at any point of the set \( \Xi(A, b) \), i.e., the united solution set to the interval linear system \( Az = b \).

Taking into account the consideration of the preceding section, the next question arises: how can we correct the construction to improve the objective function and avoid plateau-like minima sets at the zero level in the case of nonempty feasible parameter sets? This can be done in a natural way we describe in the following section.

5 Maximum Compatibility Method

To bring the results of the preceding section into correlation with known interval-related estimation techniques, it is necessary to transform the problem statement (22)–(23). We will need the following property of the mignitude (see [18, 31]): for any intervals \( u \) and \( v \),

\[
\langle u \pm v \rangle \geq \langle u \rangle - |v|.
\]
The equality instead of the non-strict inequality (25) holds true, in particular, if \( \langle u \rangle \geq |v| \) and the interval \( v \) is balanced (symmetric with respect to zero), i.e., has the form \( v = [-\tilde{v}, \tilde{v}] \) for a nonnegative \( \tilde{v} \). The short proof is as follows:

\[
\langle u \pm v \rangle = \min_{u \in U} |u \pm v| \geq \min_{u \in U} |u| - |v| = \langle u \rangle - |v|.
\]

If \( v \) is a balanced interval, then, for any \( u \in U \), there exists such \( v \in V \) that the equality \( |u \pm v| = |u| - |v| \) holds, no matter what the interval \( u \) is. Hence, the equality instead of the inequality in the above relations really takes place.

**Recognizing functional Uni.** For each index \( i = 1, 2, \ldots, m \), we can evaluate subexpressions of (23) as follows:

\[
\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \rangle = \langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid } b_i - [-1, 1] \cdot \text{rad } b_i \rangle \\
\geq \langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid } b_i \rangle - |[-1, 1] \cdot \text{rad } b_i| \\
= \langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid } b_i \rangle - \text{rad } b_i
\]

due to property (25). Notice that the equality instead of non-strict inequality is valid for the case

\[
\langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid } b_i \rangle \geq \text{rad } b_i.
\]

Since the magnitude is always nonnegative, we have

\[
\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \rangle = \max \{ \psi_i(z), 0 \},
\]

where

\[
\psi_i(z) \overset{\text{def}}{=} \langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid } b_i \rangle - \text{rad } b_i.
\]

In summary,

\[
\Phi(z) = \max_{1 \leq i \leq m} \max \{ \psi_i(z), 0 \},
\]

and, reversing the order of the two maximum operations, we get

\[
\Phi(z) = \max \{ \Psi(z), 0 \},
\]

(26)
where

$$\Psi(z) \overset{\text{def}}{=} \max_{1 \leq i \leq m} \psi_i(z) = \max_{1 \leq i \leq m} \left\{ \left\langle \sum_{j=1}^{n} a_{ij} z_j - \text{mid} b_i \right\rangle - \text{rad} b_i \right\}.$$  

One can see that $\Psi(z)$ differs only by the opposite sign from the so-called recognizing functional of the united solution set $\Xi(A, b)$ to the interval linear system $A z = b$, which is defined as

$$\text{Uni}(z) \overset{\text{def}}{=} \min_{1 \leq i \leq m} \left\{ \text{rad} b_i - \left\langle \text{mid} b_i - \sum_{j=1}^{n} a_{ij} z_j \right\rangle \right\}.$$  

It was proposed and studied in [27, 30]. Thus, the minimized functional $\Phi(z) = \max \{ \Psi(z), 0 \}$ is related to the functional $\text{Uni}(z)$ as

$$\Phi(z) = \max \{ -\text{Uni}(z), 0 \} = - \min \{ \text{Uni}(z), 0 \}.$$  

**Recognizing functional Uss.** We can perform our transformations of expression (23) for $\Phi$ in another way. For each index $i = 1, 2, \ldots, m$, there holds

$$\left\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \right\rangle$$

$$= \left\langle \sum_{j=1}^{n} (\text{mid} a_{ij} + [-1, 1] \cdot \text{rad} a_{ij}) z_j - \text{mid} b_i - [-1, 1] \cdot \text{rad} b_i \right\rangle$$

$$= \left\langle \sum_{j=1}^{n} (\text{mid} a_{ij}) z_j + \sum_{j=1}^{n} [-1, 1] (\text{rad} a_{ij}) z_j - \text{mid} b_i + [-1, 1] \cdot \text{rad} b_i \right\rangle$$

due to distributivity of multiplication with respect to addition for the common point multipliers $z_j$

$$\geq \left\langle \sum_{j=1}^{n} (\text{mid} a_{ij}) z_j - \text{mid} b_i \right\rangle - \left\lfloor \sum_{j=1}^{n} [-1, 1] (\text{rad} a_{ij}) z_j + [-1, 1] \cdot \text{rad} b_i \right\rfloor$$

due to property [25] of the magnitude

$$= \left\lfloor \sum_{j=1}^{n} (\text{mid} a_{ij}) z_j - \text{mid} b_i \right\rfloor - \sum_{j=1}^{n} (\text{rad} a_{ij}) |z_j| - \text{rad} b_i.$$
since all the intervals $[-1,1] \cdot \text{rad} \mathbf{a}_{ij} z_j$, $[-1,1] \cdot \text{rad} \mathbf{b}_i$ are symmetric with respect to zero and $(\sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j - \text{mid} \mathbf{b}_i)$ is a point, not interval. Therefore,

$$\left\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \right\rangle \geq \mid \text{mid} \mathbf{b}_i - \sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j \mid - \sum_{j=1}^{n} (\text{rad} \mathbf{a}_{ij}) |z_j| - \text{rad} \mathbf{b}_i,$$

and the equality instead of non-strict inequality is valid for the case

$$\mid \text{mid} \mathbf{b}_i - \sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j \mid \geq \sum_{j=1}^{n} (\text{rad} \mathbf{a}_{ij}) |z_j| + \text{rad} \mathbf{b}_i.$$

Since the magnitude is always nonnegative, we have

$$\left\langle \sum_{j=1}^{n} a_{ij} z_j - b_i \right\rangle = \max \{ v_i(z), 0 \},$$

where

$$v_i(z) \equiv \sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j - \text{mid} \mathbf{b}_i - \sum_{j=1}^{n} (\text{rad} \mathbf{a}_{ij}) |z_j| - \text{rad} \mathbf{b}_i.$$

In summary,

$$\Phi(z) = \max_{1 \leq i \leq m} \max \{ v_i(z), 0 \},$$

and, reversing the order of the two maximum operations, we get

$$\Phi(z) = \max \{ \Upsilon(z), 0 \},$$

(27)

where

$$\Upsilon(z) \equiv \max_{1 \leq i \leq m} \left\{ \sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j - \text{mid} \mathbf{b}_i - \sum_{j=1}^{n} (\text{rad} \mathbf{a}_{ij}) |z_j| - \text{rad} \mathbf{b}_i \right\},$$

One can see that $\Upsilon(z)$ differs only by the opposite sign from the so-called recognizing functional of the united solution set $\Xi(\mathbf{A}, \mathbf{b})$ to the interval linear system $\mathbf{A}z = \mathbf{b}$, which is defined as

$$\text{Uss}(z) = \min_{1 \leq i \leq m} \left\{ \text{rad} \mathbf{b}_i + \sum_{j=1}^{n} (\text{rad} \mathbf{a}_{ij}) |z_j| - \text{mid} \mathbf{b}_i - \sum_{j=1}^{n} (\text{mid} \mathbf{a}_{ij}) z_j \right\}.$$

It was introduced and investigated in [28, 29, 32]. Thus, the minimized functional $\Phi(z) = \max \{ \Upsilon(z), 0 \}$ is related to the functional $\text{Uss}(z)$ as

$$\Phi(z) = \max \{ -\text{Uss}(z), 0 \} = - \min \{ \text{Uss}(z), 0 \}.$$
The sense and general properties of recognizing functionals. The recognizing functionals Uni and Uss were introduced in [27, 28, 29, 30, 32] to give a numerical measure that characterizes compatibility of interval linear equation systems.

According to the result from [30], a point $z \in \mathbb{R}^n$ belongs to the united solution set $\Xi(A, b)$ if and only if
\[
\langle Az - \text{mid} b \rangle \leq \text{rad} b,
\]
where the magnitude is applied component-wise. So, the amount to which the right-hand side of the above inequality exceeds its left-hand side, i.e., the difference
\[
\text{rad} b - \langle Az - \text{mid} b \rangle,
\]
can be taken as a numerical measure of compatibility between the parameter vector $z$ and the data $A, b$. In order to construct a unified scalar characteristic, we convolve vector (28) by taking the minimum of its components. This way, the functional Uni is obtained.

The membership of a point $z \in \mathbb{R}^n$ in the united solution set $\Xi(A, b)$ to the interval linear equation system $Az = b$ is equivalent to the non-negativeness of the functional Uni in $z$:
\[
z \in \Xi(A, b) \iff \text{Uni} (z, A, b) \geq 0.
\]

Well-known Oettli-Prager inequality (see [18, 23]) may serve as a base for yet another compatibility measure. Namely, a point $z \in \mathbb{R}^n$ is known to belong to the united solution set $\Xi(A, b)$ if and only if
\[
\mid \text{mid} A \cdot z - \text{mid} b \mid \leq \text{rad} A \cdot |z| + \text{rad} b,
\]
where the magnitude (absolute value) is applied componentwise. So, the amount to which the right-hand side of the above Oettli-Prager inequality exceeds its left-hand side, i.e., the difference
\[
\text{rad} A \cdot |z| + \text{rad} b - \mid \text{mid} A \cdot z - \text{mid} b \mid,
\]
can be taken as a numerical measure of compatibility between the parameter vector $z$ and the data $A, b$. In order to construct a unified scalar characteristic, we convolve vector (29) by taking the minimum of its components. This way, the functional Uss is obtained.

The membership of a point $z \in \mathbb{R}^n$ in the united solution set $\Xi(A, b)$ to the interval linear equation system $Az = b$ is equivalent to the non-negativeness of the functional Uss in $z$:
\[
z \in \Xi(A, b) \iff \text{Uss} (z, A, b) \geq 0.
\]

Below, we give a short survey of the properties of the recognizing functionals Uni and Uss as they are presented in the works [27, 28, 29, 30, 32].
The functionals $\text{Uni}$ and $\text{Uss}$ are concave functions of $z$ in each orthant of the space $\mathbb{R}^n$. If, additionally, the matrix $A$ is such that its columns with the indices from the set $J = \{j_1, j_2, \ldots, j_r\}$, $r \leq n$, are intervals, while the rest columns are real numbers (degenerate intervals), then the functionals $\text{Uni}$ and $\text{Uss}$ are concave on each of the $2^r$ sets of the form $\{z \in \mathbb{R}^n \mid z_j \geq 0, j \in J\}$, where “$\geq$” denotes one of the relations “$\geq$” or “$\leq$”.

The functionals $\text{Uni}$ and $\text{Uss}$ are polyhedral, that is, their graphs are made up of finite numbers of hyperplane pieces (that can be seen at Fig 8). Additionally, the functional $\text{Uni}$ reaches a finite maximum with respect to $z$ over the entire space $\mathbb{R}^n$.

On the other hand, for interval linear systems with nonempty united solution set, the recognizing functionals can help distinguishing interior and boundary of the solution set, etc. In particular, if $\text{Uni}(z) > 0$ or $\text{Uss}(z) > 0$, then $z$ is a point from the topological interior of the solution set, which immediately follows from the continuity of $\text{Uni}$ and $\text{Uss}$. Under certain additional requirements, the converse is also true [28, 29]. Overall, the recognizing functionals prove to be very useful tool for examination of the “fine structure” of the solution sets.

Both functionals, $\text{Uss}$ and $\text{Uni}$, have similar properties, and each one of them is intended for its own problems. In particular, the functional $\text{Uni}$ takes into account the right-hand side of the system to a greater extent than the functional $\text{Uss}$. Hence, $\text{Uni}$ may be useful in the problems where the corresponding variables are of much importance.

The recognizing functionals may be considered as a tool to find a kind of solution to a system of interval equations when we have underestimated the uncertainty with which we know the parameters. The corresponding system of interval linear equations has no solutions, i.e., its united solution set is empty. Then one can use the vector for which the functional $\text{Uni}$ or $\text{Uss}$ attains its largest possible value as a pseudo-solution with the minimum possible incompatibility.

An example of interval linear system and its recognizing functionals.

As an example, we consider the interval linear algebraic system

\[
\begin{pmatrix}
[2, 4] & [-2, 0] \\
[-1, 1] & [2, 4]
\end{pmatrix}
\begin{pmatrix}
z_1 \\
z_2
\end{pmatrix} =
\begin{pmatrix}
[-1, 1] \\
[0, 2]
\end{pmatrix},
\] (30)

and its solution set is depicted at Fig. 7.

The pictures at Fig. 8 show graphs of the recognizing functionals $\text{Uni}$ and $\text{Uss}$ of the solution set for system (30). The difference between the two pictures is small, but it exists: at the graph of $\text{Uss}$, the maximum is a “sharp peak” of the height 1.44 attained at the point $(0.1, 0.33)^\top$, while the graph of $\text{Uni}$ has a flat maximum region at the height 1 around the argument point $(0.5, 0.5)^\top$.

For negative values, the functionals $\text{Uni}$ and $\text{Uss}$ totally coincide with each other, as was substantiated at the beginning of the section. But for nonnegative
values, each one of them shows its own “compatibility measure”, according to their constructions, for the data given by the interval linear system (30).

**Maximum Compatibility Method.** In view of the above results, the situation when the interval system has empty solution set corresponds to the case when $\Psi(z) > 0$ and $\Upsilon(z) > 0$ for all $z$, i.e., when

$$\text{Uni}(z) = - \Psi(z) < 0 \quad \text{and} \quad \text{Uss}(z) = - \Upsilon(z) < 0 \quad \text{for all } z \in \mathbb{R}^n.$$ 

Then we have

$$-\Phi(z) = \min \{ \text{Uni}(z), 0 \} = \text{Uni}(z)$$

and, at the same time,

$$-\Phi(z) = \min \{ \text{Uss}(z), 0 \} = \text{Uss}(z).$$

Hence, minimizing $\Phi(z)$ in our problem (22)–(23) is equivalent to unconstrained maximization of $\text{Uni}(z)$ or $\text{Uss}(z)$.

The approach to data fitting and parameter estimation based on maximization of the recognizing functional for the solution set is called the **Maximum Compatibility Method** (the name “maximum consistency method” was also used in the previous works [27, 32]). As an estimate of parameters, we take the argument of the recognizing functional where its maximum is attained. The term “maximum compatibility” is justified by the fact that the recognizing functionals, both $\text{Uni}$ and $\text{Uss}$, show, as was demonstrated earlier, the degree of compatibility between the data $A, b$ and parameters $z_1, z_2, \ldots, z_n$ of the regression line. If the maximum of the recognizing functional is nonnegative, then our estimate is compatible with the input data in the sense of the definition from Section 3.
Figure 8: Graphs of the recognizing functionals Uni and Uss for system (30).
If the maximum of the recognizing functional is negative, no compatibility can be achieved between any parameter estimate and the data, but the point we have found is the best one, since it provides the minimum possible value of “incompatibility”. In the limit case of non-interval (point) data, maximization of any recognizing functional is equivalent to minimizing the Chebyshev norm (maximum-norm) of the defect of the equation system (20) (see [32]). The Maximum Compatibility Method thus turns into the so-called “Chebyshev data smoothing” which is successfully applied in data processing.

The idea of the Maximum Compatibility Method comes from the interval analysis, while the idea of minimizing the functional $\Phi(z)$ in (22)–(23) comes from a natural probabilistic approach, the Maximum Likelihood Method. In the case of empty feasible parameter set, the estimates produced by these two methods coincide. The Maximum Compatibility Method, thereby, acquires a probabilistic justification.

To improve the situation with the zero-level minimum plateau in the straight line fitting problem (22)–(23), it makes sense to change the objective function $\Phi$ within the nonempty solution set $\Xi(A, b)$. We can take either $\Psi$ instead of $\Phi = \max\{\Psi, 0\}$, or $\Upsilon$ instead of $\Phi = \max\{\Upsilon, 0\}$, thus removing the cutoff with zero in both cases. This will delete the zero plateau at the minimum of $\Phi$ for nonempty feasible parameter set. On the other hand, this idea has already been implemented in the Maximum Compatibility Method, where one maximizes the recognizing functionals $\text{Uni} = -\Psi$ or $\text{Uss} = -\Upsilon$. Moreover, the Maximum Compatibility Method is a uniform numerical procedure in which an automatic switching takes place, depending on a specific situation, between the two dissimilar parts of the data fitting problem under interval uncertainty that correspond to empty and nonempty feasible parameter set.

6 Practical example

As a practical example, we consider an electrochemistry problem. This is the problem of determining the parameters of the temperature dependency of electrolyte ionic conductivity based on the unique experimental data from [34]. Previously, it has been treated by interval methods in [10]. The problem setting is not the most general, but the function that we fit to data is nonlinear, which helps us illustrate some important concepts of the interval data fitting.

**Problem statement.** The functional dependency under study has the form

$$S(y) = V \exp(\alpha y) + BG,$$

where $S$ is the electrolyte conductivity, $y$ is the independent variable, and the quantities $\alpha$, $V$, $BG$ are to be determined from the measurements. $BG$ is an entire identifier that means the so-called background component. The independent variable $y$ is expressed from the absolute temperature $T$ of melted electrolyte as $(T/K)^{-1}$ with a known constant $K$. 
We have to process a sample of eight measurements, performed at the values \( y_i, i = 1, 2, \ldots, 8 \), of the independent variable \( y \) that are equal to

\[
0.002481, 0.002544, 0.002583, 0.002658, 0.002785, 0.002915, 0.003002, 0.003124.
\]

Electrolyte conductivity values in decimal logarithmic scale are

\[
-1.339, -1.473, -1.561, -1.715, -1.994, -2.149, -2.205, -2.275,
\]

while in natural scale (obtained from (33) by exponentiation with the base 10) they correspond to the conductivities \( S_i = S(y_i), i = 1, 2, \ldots, 8 \), equal to

\[
0.0458142, 0.0336512, 0.0274789, 0.0192752, 0.0101391, 0.0070958, 0.0062373, 0.0053088.
\]

We have to find values of \( \alpha, V \), and \( BG \) for which the functional dependency of the form (31) best fits the experimental data (32)–(34). Due to small number of the measurements in the sample, using the statistical methods based on the theory of probability is unfounded and inadequate in the above problem.

It is worthwhile to note that, in the problem statement, the parameters \( \alpha, V \), and \( BG \) from the function (31) are not completely equivalent to each other. The main quantities that interest electrochemists are \( \alpha \) and, to a smaller extent, \( BG \). It can be said that \( BG \) plays the role of an auxiliary background constant that reflects individual specificity of the experiments. Its value is necessary to bound a part of the domain of function (31) at which the functional dependency really takes place, i.e., in essence, to estimate the maximal argument for which the experiments should be performed. For larger arguments of function (31), we get to that part of its domain where the function values are too small and do not have physical meaning.

Relying on the above understanding of the problem, the electrochemists introduce “compensated conductivity” \( \tilde{S}(y) \) obtained by subtracting the background value \( BG \) from the measurement data:

\[
\tilde{S}(y) = S(y) - BG.
\]

In its turn, the compensated conductivity produces a “compensated sample” \( \tilde{S} \) of the measurements of \( S(y) \) defined as

\[
\tilde{S}_i = S_i - BG, \quad i = 1, 2, \ldots, 8.
\]

Then the functional dependency (31) gets a simpler form

\[
\tilde{S}(y) = V \exp(\alpha y),
\]

which, after taking natural logarithm, becomes linear with respect to the input variable:

\[
\ln \tilde{S}(y) = \ln V + \alpha y.
\]
Physical and chemical reasons dictate such a value of the background component \( BG \) that the functional dependency \( (35) - (36) \) is satisfied “most accurately” for the data \( (32) - (34) \).

The independent variable \( y \) is assumed to be known exactly in the experiments under study, while the measurements \( S_i \) are subject to errors about which we only know that their upper bound is a certain value \( E \). Therefore, instead of \( (34) \), we really have a family of intervals

\[
S_i = [S_i - E, S_i + E], \quad i = 1, 2, \ldots, 8, \tag{37}
\]

with the midpoints from \( (34) \) and radii \( E \), to which the true values of the conductivity belong. These intervals will be called measurement uncertainty intervals. Taking into account the capability of the measuring devices and the experimental conditions, the value \( E = 0.002 \) has been assigned as a safe upper bound on the measurement error in \( [10] \), although the actual error is, as a rule, much smaller. De facto, after the background component \( BG \) is fixed, we have to process the interval compensated sample

\[
\tilde{S}_i = [\tilde{S}_i - E, \tilde{S}_i + E], \quad i = 1, 2, \ldots, 8, \tag{38}
\]

based on which \( \alpha \) and \( V \) are to be determined.

Due to the physical meaning of the problem, the conductivity \( S(y) \) cannot be negative or even zero. That imposes natural constraints on the value of the background component \( BG \) and actual error margin \( E \), i.e., on the radius of the measurement uncertainty intervals. If a measurement uncertainty interval contains negative values, it is senseless, and the corresponding interval should be somehow corrected.

**The ideas behind our solution.** The above problem is a data fitting problem with a nonlinear functional dependency. So, we cannot directly avail ourselves of the results on straight line fitting from Sections 4–5. A natural idea is to reformulate the Maximum Compatibility Method of Section 5 in an equivalent manner that would be applicable to our nonlinear situation. Our solution of the problem is, in essence, based on the ideas first stated in \([21, 37, 38]\) and then developed in the works \([27, 28, 29, 32]\).

Notice that the specific form of the expressions that determine the recognizing functionals \( U_i \) and \( Uss \) allows us to easily predict how their values change after varying the matrix \( A \) and right-hand side \( b \) of the interval linear algebraic systems \( Ax = b \). In both expressions

\[
\min_{1 \leq i \leq m} \left\{ \text{rad } b_i - \left( \text{mid } b_i - \sum_{j=1}^{n} a_{ij} x_j \right) \right\}
\]

and

\[
\min_{1 \leq i \leq m} \left\{ \text{rad } b_i + \sum_{j=1}^{n} (\text{rad } a_{ij}) |x_j| - \left| \text{mid } b_i - \sum_{j=1}^{n} (\text{mid } a_{ij}) x_j \right| \right\},
\]
Figure 9: Inflating the uncertainty boxes inevitably results in compatibility of the parameters and data.

the quantities $\text{rad} \ b_i$, $i = 1, 2, \ldots, m$, occur as add-ons in every subexpression standing under “min” operation. Therefore, if all $\text{rad} \ b_i$’s simultaneously increase or decrease by equal values, then the general minima increase or decrease by the same value.

Uniform increase of the radii of the right-hand sides $b_i$ by $C$, $C \geq 0$, is equivalent to adding the vector $Ce$ to $b$, where $e = ([−1,1],\ldots,[−1,1])^T$. Hence, for the interval system $Ax = b + Ce$ with the widened right-hand side, there holds

$$
\text{Uni} \ (x, A, b + Ce) = \text{Uni} \ (x, A, b) + C,
$$

$$
\text{Uss} \ (x, A, b + Ce) = \text{Uss} \ (x, A, b) + C,
$$

and, as a consequence,

$$
\max_{x \in \mathbb{R}^n} \text{Uni} \ (x, A, b + Ce) = \max_{x \in \mathbb{R}^n} \text{Uni} \ (x, A, b) + C, \quad (39)
$$

$$
\max_{x \in \mathbb{R}^n} \text{Uss} \ (x, A, b + Ce) = \max_{x \in \mathbb{R}^n} \text{Uss} \ (x, A, b) + C. \quad (40)
$$

If the solution set $\Xi(A,b)$ is empty, so that

$$
M = \max_{x \in \mathbb{R}^n} \text{Uni} \ (x, A, b) = \max_{x \in \mathbb{R}^n} \text{Uss} \ (x, A, b) < 0
$$

(it makes sense to remind that the negative values of the functionals Uni and Uss coincide), then increasing radii of all the right-hand side components by $C \geq |M|$ makes the solution set non-empty: then both $\max_{x \in \mathbb{R}^n} \text{Uni} \ (x, A, b + Ce)$ and $\max_{x \in \mathbb{R}^n} \text{Uss} \ (x, A, b + Ce)$ becomes non-negative due to (39)–(40). We can use


such information for correcting, in a necessary sense, the interval linear system, i.e., the input data for the parameter identification problem.

On the other hand, a practical interpretation of the Maximum Compatibility Method ensues from (39)–(40): the value of the argument providing negative max Uni or max Uss is the first point that appears in the solution set after uniform (with respect to the midpoint) widening of the right-hand side vector.

Yet another interpretation of the maximum compatibility method may be as follows: the arguments of negative max Uni and max Uss gives us parameters of such a regression line that should be widened in the smallest possible amount to result in a “regression strip” intersecting all the data boxes (see Fig. 10).

![Figure 10: Regression strip (instead of thin line) as a solution to data fitting problem under interval uncertainty.](image)

The above properties of the maximum compatibility estimate can lay down a basis for an approach that may be applied to nonlinear estimation problems. The main idea is to vary the amount of data uncertainty expressed in terms of the width of intervals and investigate the compatibility of the resulting sample. The minimum width of the uncertainty intervals that makes the sample compatible characterizes the ineradicable dispersion of the data with respect to the parametric function family to be fitted. The estimate of the parameter should be chosen so that it minimizes such dispersion, which is completely similar to parameter estimation in probabilistic models.

**Our plan of action.** Addressing our problem, we can note that the samples of the point measurement data (33)–(34) are usually not compatible with any of parameter sets of the functional dependencies (31) or (36) in the sense of definition from Section 3. However, as follows from the above piece of theory, widening the data uncertainty intervals (33)–(34) will always result in nonempty feasible parameter set, while the interval sample obtained gains compatibility
with a parameter set of the model under study. The smallest amount of the uniform inflation of the uncertainty intervals with respect to the specified mid-points required for the sample to become compatible is an objective measure of the data dispersion within the sample for a given functional dependency. In the linear case we have studied in Sections 4 and 5 this smallest amount of widening is equal to the absolute value of the maximum of the recognizing functional. In the general nonlinear problem we solve in the current section, such a simple relationship is not valid any more, but the smallest amount of the inflation of point data \( \hat{S} \), in natural scale, necessary for the compatibility to be attained, can be taken, for given parameter values, as a measure of their compatibility with the data. This way, we implicitly estimate the minimal level of the measurement errors present in the processed sample.

To inquire into compatibility of the interval compensated sample \( \hat{S} \), we transform, by taking the logarithm, the constructed functional dependency to the linear form \( W = \ln \hat{S} \). In doing that, we have to compute \( \ln \hat{S}_i \), \( i = 1, 2, \ldots, 8 \), i.e., logarithms of the interval data \( \hat{S} \). According to the sense of the problem under solution, these are interval of values of logarithms of the numbers from intervals \( \hat{S} \), so that they can be easily computed based on monotonicity of the logarithm. Further, the compatibility of the logarithmed sample with respect to the linear dependency \( W \) is equivalent to compatibility (solvability) of the interval linear system of equations

\[
W u = \ln \hat{S},
\]

in which \( W \) is an \( 8 \times 2 \)-matrix of the form

\[
W = \begin{pmatrix}
1 & y_1 \\
1 & y_2 \\
\vdots & \vdots \\
1 & y_8
\end{pmatrix},
\]

\( u = (\ln V, \alpha)^\top \), \( \ln \hat{S} = (\ln \hat{S}_1, \ldots, \ln \hat{S}_8)^\top \). The specific form of the matrix \( W \) in the interval linear system \( W \) is explained by the fact that, in our data fitting problem, the linear function \( \ln \hat{S} \) has a constant term. Finally, to determine whether system \( W \) is compatible or not, we find unconstrained maximum of the recognizing functional \( Uss(W, \ln \hat{S}, u) \) over the variable \( u \) and compare it with zero. As far as the matrix \( W \) is point (non-interval), the recognizing functional \( Uss \) is globally concave, and we can use the computer code \texttt{lintregs} for the maximization of \( Uss \).

**Computational technology and results.** To describe formally our computational procedure, we introduce a function \( E_{\text{min}}(BG) \) that assigns, to every value of the background component \( BG \), the smallest radius \( E \) of the measurement uncertainty intervals for which the interval compensated sample \( \hat{S} \) is compatible. \( E_{\text{min}}(BG) \) is thus the function producing the data dispersion for
the compensated sample with the background $BG$. Therefore, our problem can be solved by minimizing $E_{\text{min}}(BG)$ over $BG$ from its domain, i.e., from the set of its meaningful values.

It follows from the above paragraph that the value of the function $E_{\text{min}}(BG)$ is nothing but a (unique) root of the equation

$$\max_{u \in \mathbb{R}^2} \text{Uss} \left( W, \ln(\tilde{S} + \theta e), u \right) = 0 \quad (42)$$

with respect to the real unknown variable $\theta$. The matrix $W$ and the vector $u$ have been defined earlier, $e = ([−1, 1], \ldots, [−1, 1])^\top$, and the logarithm $\ln$ is applied to the vector $(\tilde{S} + \theta e)$ in componentwise manner. We solved equation (42) by bisection (dichotomy) method, taking $[0, 2E] = [0, 0.004]$ as the initial interval where the root is localized.

The main stage of the solution of our problem is minimization of the function $E_{\text{min}}(BG)$ over $BG \in [0, 0.005]$. Algorithmically, it can be organized as the one-dimensional direct search, for instance, using “golden section” search or any other analogous approach (see [1] and the other textbooks on numerical optimization). We got that, for the background variable $BG = 0.003757$, the smallest radius of the intervals of the compensated sample $\tilde{S}$ which is necessary for its compatibility is equal to merely $E^*_{\text{min}} = 0.0006197$ being the minimal one among all $E_{\text{min}}$ for any other values of the background variable $BG$. Then $\alpha^* = −5945.7$ and $\ln V^* = 11.597$, so that the sought-for functional dependency (31), constructed in accordance to our “smallest data dispersion” criterion, has the form

$$S(y) = e^{11.597} \exp(−5945.7 y) + 0.003757$$

$$= 108786 \cdot \exp(−5945.7 y) + 0.003757.$$

7 Remaining open problems

To conclude our spacious paper, it makes sense to formulate some open questions. Answering them would be further advance in non-probabilistic statistics and interval data analysis techniques.

What are the asymptotic properties of the maximum compatibility estimates? In the probabilistic situation, when we use traditional statistical procedures, it is well-known that, under certain reasonable conditions, the resulting estimates tend to the actual values of the corresponding quantities as the number of measurements increases. Moreover, we can also estimate the rate at which the estimates tend to the actual values. Intuitively, it looks like a similar asymptotic property should hold for interval methods as well.

This is definitely true in the simplest case when we only have one scalar parameter $p = p_1$, and the measured quantity $y$ does not depend on the quantities $x_i$ at all, i.e., $f(x, p) = p_1$. Then each measurement result has the form $y^{(i)} = p + \Delta y^{(i)}$, $i = 1, 2, \ldots, m$, where measurement errors $\Delta y^{(i)}$ are
independent implementations of a probability distribution located on the interval $[-\Delta, \Delta]$. Consequently, the actual value $p$ is located in every interval $[y^{(i)} - \Delta, y^{(i)} + \Delta]$, $i = 1, 2, \ldots, m$. After $m$ measurements, we can thus conclude that $p$ is contained in the intersection of the corresponding $m$ intervals, i.e., in the interval

$$\bigcap_{i=1}^{m} [y^{(i)} - \Delta, y^{(i)} + \Delta] = \left[ \max_{1 \leq i \leq m} y^{(i)} - \Delta, \min_{1 \leq i \leq m} y^{(i)} + \Delta \right]$$

(43)

$$= p + \left[ \max_{1 \leq i \leq m} \Delta y^{(i)} - \Delta, \min_{1 \leq i \leq m} \Delta y^{(i)} + \Delta \right].$$

In the non-degenerate case, i.e., unless the error is located with probability 1 in a proper subinterval of the interval $[-\Delta, \Delta]$, it can be proven that the intersection (43) tends to 0 as $m$ increases [36]. We really get an asymptotic convergence, and the corresponding error turns out to decrease as $O(1/m)$ (see details in [36] too). Moreover, the convergence obtained is asymptotically faster than, e.g., in the Gaussian probabilistic case.

Indeed, in the probabilistic case, when the measurement error is normally distributed with zero mean, some standard deviation $\sigma$ and variance $V = \sigma^2$, the optimal way to estimate the actual value $p$ based on the measurement results $y^{(1)}, \ldots, y^{(m)}$ is known to compute the arithmetic average

$$\bar{p} = \frac{y^{(1)} + \ldots + y^{(m)}}{m}.$$

It is easy to compute the standard deviation $\sigma_{\bar{p}}$ of this arithmetic average. Namely, the variance $V_s$ of the sum

$$s \overset{\text{def}}{=} y^{(1)} + \ldots + y^{(m)}$$

of $m$ independent random variables is equal to the sum of their variances, i.e., to $V_s = Vm = \sigma^2m$. Therefore, the standard deviation $\sigma_s = \sqrt{V_s}$ of this sum is equal to $\sigma\sqrt{m}$. When we divide a random variable by a positive constant $m$, its standard deviation divides by the same constant. Thus, for $\bar{p} = s/m$, the standard deviation $\sigma_{\bar{p}}$ is equal to

$$\sigma_{\bar{p}} = \frac{\sigma_s}{m} = \frac{\sigma\sqrt{m}}{m} = \frac{\sigma}{\sqrt{m}}.$$

So, for the Gaussian probabilistic uncertainty, the inaccuracy of the resulting estimate decreases as $O(1/\sqrt{m})$, which is much slower than $O(1/m)$ for the interval uncertainty [36].

It is desirable to extend the result on asymptotic convergence from [36] to a general case, in which there can be several parameters $p_j$, the functional dependency $f(x, p)$ is not so simple, and the values $x^{(i)}$ can change from one measurement to another.
What if we have partial information about the probabilities? In this paper, we consider only two extreme situations: the probabilistic situation when we know the probability distributions of all the measurement errors, and the interval situation, in which the only information that we have about the measurement error is the upper bound on its absolute value.

In practice, in addition to the upper bound, we often have some partial information about the probability distribution of the measurement error too; see, e.g., [17]. For example, we may have bounds on some moments of this distribution, or we may know bounds on the values of the corresponding cumulative distribution function, etc. In such situations, we can also apply the maximum entropy approach to specify the shape of the probability distribution (as in Section 2), but now, when we have additional information about the distribution, this method may lead to non-uniform distributions, as distinct from the conclusion [11]. It is desirable to translate this general idea into efficient techniques that would thus generalize the Maximum Compatibility Method to the wider applicability scope.

In this case, we can raise a similar question about the asymptotic properties of the resulting estimates.

References


[12] lintregr.m — a free MATLAB code implementing the Maximum Compatibility Method for interval data fitting. Available at [http://www.nsc.ru/interval/Programing/MCodes/lintregr.m](http://www.nsc.ru/interval/Programing/MCodes/lintregr.m)


