# eistoolbox

for

# MathWorks® MATLAB

A toolbox for batch fitting of Electrochemical Impedance Spectroscopy data to equivalent circuit models

# User Guide

Juan J. Montero-Rodríguez

# **Contents**

1	Introduction							
2	The	Theory						
	2.1	Imped	lance Spectroscopy	5				
	2.2	2 Kramers-Kronig Transforms						
	2.3	Fitting	g algorithms	6				
		2.3.1	Levenberg-Marquardt	6				
3	Qui	ck Star	t Guide	7				
	3.1	Graph	nical User Interface	8				
		3.1.1	Main Window	8				
		3.1.2	Results Window	8				
		3.1.3	Correlations Window	9				
4	<b>Equivalent Circuits</b>							
		4.0.1	Circuit Elements	11				
		4.0.2	Circuit String Syntax	11				
		4.0.3	Circuit Files (.ckt)	11				
5	Fitti	Fitting Algorithms 13						
	5.1	Weighting						
	5.2	Imple	mented algorithms	13				
		5.2.1	fminsearchbnd	13				
	5.3	Other	algorithms	14				
6	Stat	istics		15				
	6.1	Linear	r regressions	15				
		6.1.1	Real of fitted vs Real of measured	15				
		6.1.2	Imag of fitted vs Imag of measured	15				
		6.1.3	MAG of fitted vs MAG of measured	15				
	6.2	Good	ness of fit	15				
		621	Chi-square	15				

	6.2	6.2.2 Test F	
7		Error estimates for individual parameters	15 16
	7.1	Zfit	16
	7.2	fminsearchbnd	16

## Introduction

**eistoolbox** is a toolbox for MATLAB® used for batch fitting Electrochemical Impedance Spectroscopy (EIS) data to equivalent circuits models.

In EIS the **impedance** of a sample is observed and recorded at different frequencies, and the impedance is stored as a Bode diagram, with magnitude and phase (polar format). Results can also be stored as real and imaginary parts (rectangular format) and plotted as a Nyquist diagram.

This impedance is commonly fitted to circuit models, selected depending on the physical characteristics of the measured system. The result is a list of **circuit elements** such as resistors, capacitors and inductors, and a list of **parameter values**.

The problem is how to calculate these parameters. This is often achieved using **nonlinear optimization**, and some of the most famous algorithms for fitting EIS data are based on the Complex Nonlinear Least Squares (CNLS) algorithm such as the Levenberg-Marquardt. However there are many algorithms for this end, and they are difficult to implement as they require advanced mathematics.

The present **eistoolbox** is a collection of algorithms for nonlinear optimization, and a graphical user interface for batch fitting of multiple EIS curves.

# Theory

#### **Impedance Spectroscopy** 2.1

Electrochemical Impedance Spectroscopy (EIS) measures the complex impedance of a sample as a function of the frequency. The experimental results are stored in two possible formats: polar coordinates (magnitude and phase) or rectangular coordinates (real and imaginary).

$$Z(f) = R + jX (2.1)$$

where R is the resistance and X is the reactance of the sample.

The real part of the impedance is proportional to the resistivity, and the imaginary part is proportional to the permittivity. Both parameters can be calculated directly from the measurements, considering the exact geometry of the electrodes and measurement setup. For parallel plate electrodes, the following equations apply:

$$R = \frac{\rho L}{A} \tag{2.2}$$

$$R = \frac{\rho L}{A}$$

$$C = \frac{\epsilon A}{D}$$
(2.2)

The capacitive reactance is given by

$$X_C = \frac{1}{2\pi f C} \tag{2.4}$$

Substituting (2.3) and (2.4) into (2.1) results in the following equation, which describes the impedance in terms of the resistivity and permittivity of the sample between parallel electrodes:

$$Z(f) = \frac{\rho L}{A} + \frac{1}{2\pi f} \frac{D}{\epsilon A} \tag{2.5}$$

Impedance Spectroscopy is also referred as Dielectric Spectroscopy, because it gives information about the dielectric properties of the measured sample.

CHAPTER 2. THEORY 6

## 2.2 Kramers-Kronig Transforms

Experimental data can be validated using the Kramers-Kronig Transforms.

### 2.3 Fitting algorithms

### 2.3.1 Levenberg-Marquardt

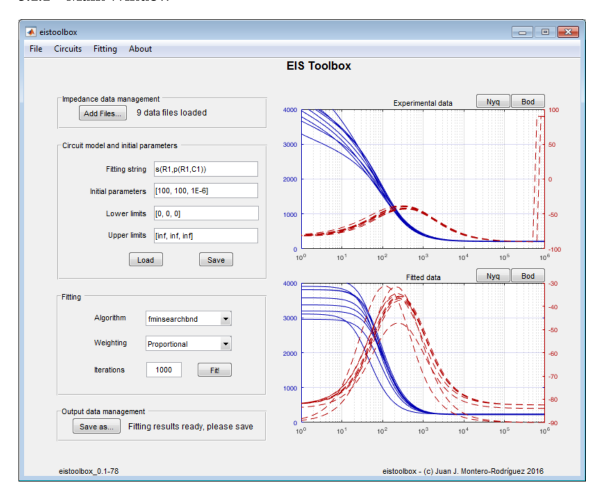
# **Quick Start Guide**

Steps for batch fitting impedance data curves:

- 1. Add files using the "Add file..." button (supported .CSV or .DTA)
- 2. Write the circuit string and fitting parameters (circuit string formatting, see Chapter 3)
- 3. Click the "Fit" button
- 4. Save the results using the "Save..." button

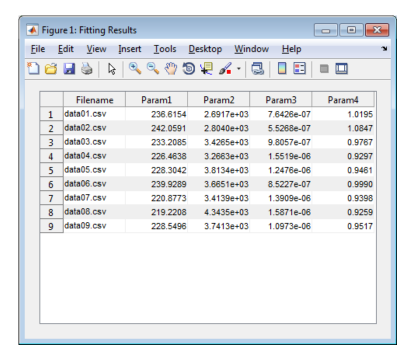
### 3.1 Graphical User Interface

#### 3.1.1 Main Window



#### 3.1.2 Results Window

The output parameters after the fitting are displayed as a table, in the same order as they appear in the circuit string.



#### 3.1.3 Correlations Window

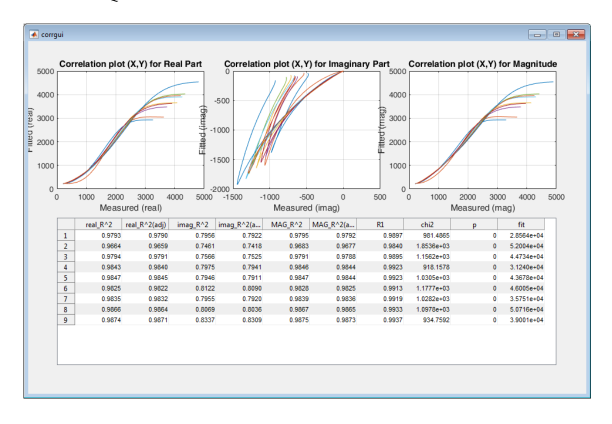
This window is useful to validate the fitting results.

There are three correlations implemented: real, imaginary and magnitude.

- The real correlation coefficient (rsq\_real) is obtained by linear regression from the plot of the fitted vs. measured real part of the impedance.
- The imaginary correlation coefficient (rsq\_imag) is obtained by linear regression from the plot of the fitted vs. measured imaginary part of the impedance.
- The magnitude correlation coefficient (rsq\_MAG) is obtained by linear regression from the plot of the fitted vs. measured magnitude of the impedance.

The plots themselves are presented as individual figures. For a perfect fit, the three plots are straight lines where y(x)=x.

Additionally, the fitting results are compared with the original data files by using the Pearson's Chisquare Test of Goodness-of-Fit. This information includes the following parameters: R, chi2, p, fit. These parameters are further explained in Chapter 5.



# **Equivalent Circuits**

Experimental data is fitted to equivalent circuit models. The models are designed to describe the interfaces, chemical processes and boundaries of the measured setup.

#### 4.0.1 Circuit Elements

The elements present in the software are described in Table 4.1:

Table 4.1: Equivalent circuit elements and their MATLAB implementation.

Symbol	Element	Equation	MATLAB expression
R1	Resistor	Z(f) = R	z=p*ones(size(f))
C1	Capacitor	$Z(f) = 1/j2\pi fC$	z=1./(1i*2*pi*f*p)
L1	Inductor	$Z(f) = j2\pi f L$	z=1i*2*pi*f*p
E2	Constant Phase Element	$Z(f) = 1/p_1(j2\pi f)^{p_2}$	$z=1./(p(1)*(1i*2*pi*f).^p(2))$

The Warburg element can be obtained with a Constant-Phase Element by setting  $p_2 = 0.5$ 

#### 4.0.2 Circuit String Syntax

Circuits can be built using series and parallel combinations of the elements in Table 4.1, using the series and parallel operators s() and p(). These operators can contain any number of elements, separated by commas.

The number next to the element letter is the number of free parameters for this element. For a capacitor (C1) the only free parameter is the capacitance. For the constant-phase element (E2) the free parameters are p1 and p2.

**Common mistake:** Do not write the circuit elements as s(R1,R2,R3,C4...). The elements cannot be written as labels. Instead, write s(R1,R1,R1,C1...).

#### 4.0.3 Circuit Files (.ckt)

The circuit string, initial parameters and boundary conditions can be stored and loaded from a circuit file with the .ckt extension. This file is read by MATLAB line-to-line. The file should include only four lines with the following content:

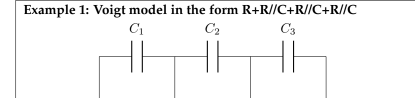
Line 1: circuit string

Line 2: initial parameters, sorted in the same order as they appear in the circuit string

Line 3: lower boundary conditions (LB)

Line 4: upper boundary conditions (UB)

Check the folder 'examples\_circuits' for more examples.



 $R_2$ 

#### Recommended circuit file

 $R_S$ 

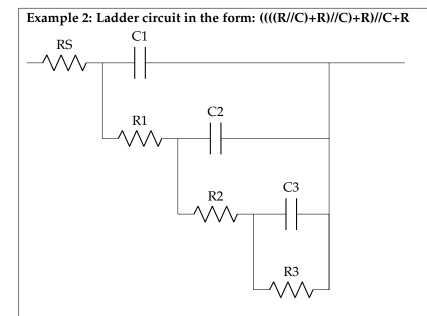
s(R1,p(R1,C1),p(R1,C1),p(R1,C1)) [100,100,1e-6,100,1e-6,100,1e-6] [0,0,0,0,0,0,0] [inf,inf,inf,inf,inf,inf]

 $R_1$ 

#### Notes

The elements appear in the circuit string in the following order: RS,R1,C1,R2,C2,R3,C3 The initial parameters, LB and UB have the exact same order

 $R_3$ 



#### Recommended circuit file

s(p(s(p(s(p(R1,C1),R1),C1),R1),C1),R1) [100,1e-6,100,1e-6,100,1e-6,100] [0,0,0,0,0,0,0] [inf,inf,inf,inf,inf,inf]

#### Notes

The elements appear in the circuit string in the following order: R3,C3,R2,C2,R1,C1,RS The initial parameters, LB and UB have the exact same order

# **Fitting Algorithms**

The fitting algorithms of this toolbox reduce the following distance function:

```
function dist=distance(param)
    ymod=feval(handlecomputecircuit, param, circuitstring, freq);
    if isequal('fitNP', fitstring)
        dist=sum(sum((ymod-zrzi).^2));
    else
        dist=sum(sum(((ymod-zrzi)./zrzi).^2));
    end
end
```

### 5.1 Weighting

The weighting type describes how the variables are compared. Currently this toolbox implements **non-proportional** and **proportional** fitting.

For **non-proportional fitting** the algorithm minimizes the squared difference between fitted (observed) and measured (expected) data.

$$d = \sum (o - e)^2$$

For **proportional fitting** the algorithm calculates the squared difference between fitted (observed) and measured (expected) data, divided by the measured (expected) data, in a way similar to the Pearson's chi-square test:

$$d = \chi^2 = \sum \frac{(o-e)^2}{e}$$

If the experimental impedance curve matches exactly the simulated curve, the distance function would have a value of zero.

### 5.2 Implemented algorithms

#### 5.2.1 fminsearchbnd

To minimize the distance function, the toolbox uses the **fminsearchbnd** algorithm.

### 5.3 Other algorithms

The following optimization algorithms are not yet implemented in this toolbox. These are some of the most used algorithms for fitting EIS data to equivalent circuit models.

- Levenberg-Marquardt
- Nelder-Mead
- BFGS
- Powell

## **Statistics**

The overall quality of the fitting can be determined by comparing the original measured data file (expected values) to the simulated values using the fit results (observed values).

Currently the software implements two methods for this comparison: linear regression and Pearson's chi-square test of goodness-of-fit.

### 6.1 Linear regressions

- 6.1.1 Real of fitted vs Real of measured
- 6.1.2 Imag of fitted vs Imag of measured
- 6.1.3 MAG of fitted vs MAG of measured
- 6.2 Goodness of fit
- 6.2.1 Chi-square

$$\chi^2 = \sum_{i}^{n} \frac{(Observed_i - Expected_i)^2}{Expected_i}$$

Observed= fitted data

Expected= measured data

#### 6.2.2 Test F

ToDo. Check Electrochemical Impedance Spectroscopy and its Applications by Adnrzej Lasia.

### 6.3 Error estimates for individual parameters

ToDo

## Licenses for included software

#### 7.1 **Z**fit

The original file was released in 2005 and it is available here:

https://de.mathworks.com/matlabcentral/fileexchange/19460-zfit

Copyright (c) 2005, Jean-Luc Dellis All rights reserved.

Redistribution and use in source and binary forms, with or without modification, are permitted provided that the following conditions are  $^{\text{met}}$ .

\* Redistributions of source code must retain the above copyright notice, this list of conditions and the following disclaimer.
\* Redistributions in binary form must reproduce the above copyright notice, this list of conditions and the following disclaimer in the documentation and/or other materials provided with the distribution

THIS SOFTWARE IS PROVIDED BY THE COPYRIGHT HOLDERS AND CONTRIBUTORS "AS IS" AND ANY EXPRESS OR IMPLIED WARRANTIES, INCLUDING, BUT NOT LIMITED TO, THE IMPLIED WARRANTIES OF MERCHANTABILITY AND FITNESS FOR A PARTICULAR PURPOSE ARE DISCLAIMED. IN NO EVENT SHALL THE COPYRIGHT OWNER OR CONTRIBUTORS BE LIABLE FOR ANY DIRECT, INDIRECT, INCIDENTAL, SPECIAL, EXEMPLARY, OR CONSEQUENTIAL DAMAGES (INCLUDING, BUT NOT LIMITED TO, PROCUREMENT OF SUBSTITUTE GOODS OR SERVICES; LOSS OF USE, DATA, OR PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY OF LIABILITY, WHETHER IN CONTRACT, STRICT LIABILITY, OR TORT (INCLUDING NEGLIGENCE OR OTHERWISE) ARISING IN ANY WAY OUT OF THE USE OF THIS SOFTWARE, EVEN IF ADVISED OF THE POSSIBILITY OF SUCH DAMAGE.

#### 7.2 fminsearchbnd

This function was written by John D'Errico and published on MathWorks MATLAB Central under an open-source license. The original file can be downloaded at https://de.mathworks.com/matlabcentral/fileexchange/8277-fminsearchbnd--fminsearchcon.

The function is based on fminsearch and includes the possibility of using boundary conditions, such as the lower and upper limits for the individual circuit parameters.