# Tuning of autoISF settings for Full Closed Loop aided by the Emulator V.2.8 for 3.0.1

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Please note that with autoISF you are in an early-dev. environment, where the user interface is **not optimized for safety** of users who stray away from intended ways to use. Good safety features exist, but these are only as good as the development-oriented user understands and implements them. This is not a medical product, refer to disclaimer in section 0



Available related case studies:

Based on older autoISF and older

Emulator versions, examples from

emulator use can be found in case

there), and case study 8.2

study 6.2, in case study 4.1 (last pages

Case study 10.1:

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10 10.1 Installing the Emulator on your PC

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10.3.3.4 .pdf chart

Rather than elaborating further, what to best do in data analysis, we should, over time, add case studies.

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You can set up and tune the system for full closed loop as described in previous sections. Doing this by frequently analyzing screenshots that must be taken in real-time of the AAPS SMB tab is

36 tedious, however.

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More elegant and precise tuning can be done with a special evaluation software for the AAPS

39 logfiles, by using the **Emulator**. In the Emulator, you can see in tabular and graphical form, which autoISF component, and other settings, contributed to SMB values that determined the glucose curve.

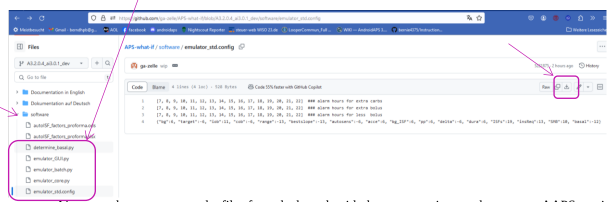
Note that iOS based variants of autoISF (Trio or iAPS) can not use the Emulator. Refer to section 11.3.

## 10.1 Installation of the emulator on your PC

## 10.1.1 Downloads

• First download **QPython 3L** onto your PC (from Google Playstore).

• Then download from: <a href="https://github.com/ga-zelle/APS-what-if/blob/A3.2.0.4\_ai3.0.1\_dev/software">https://github.com/ga-zelle/APS-what-if/blob/A3.2.0.4\_ai3.0.1\_dev/software</a>: each of these 5 py resp. config. files. To do this, you must press, for each one, here



Always make sure you use the files from the branch with the same version number as your AAPS version (in the example above: These files will work with AAPS dev version 3.2.0.4 with autoISF version 3.0.1); (the dev connotation is temporary). Always keep your AAPS x autoISF and also the emulator related files up-to-date!

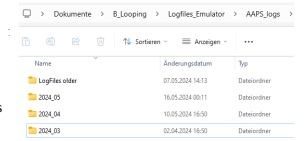
#### **10.1.2** Create your PC folder structure

Retrieve these 5 downloaded files on your PC (list of recent downloads), and shift them into a folder in

which, or neighboring to which, you also keep your logfiles.

(These you must copy-in about once a week from your phone )

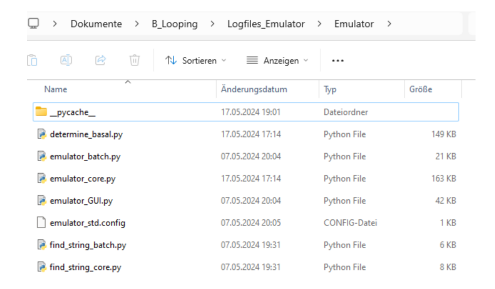
My folder structure for Logfiles and Emulation on the PC has (three) monthly folders that I most likely will look at, plus one folder with data from previous months and years.



It is advisable to additionally store a pdf from **Nightscout Reporter** in the file for every month. From it, you can much easier find which days and times are of high interest to analyze with the Emulator.

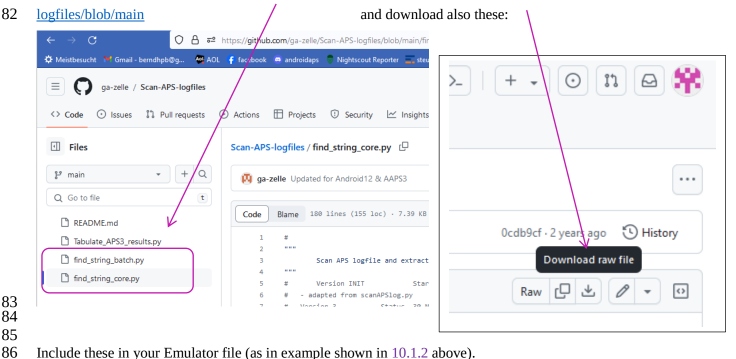
Always factor in the time difference between your AAPS phone and the "Z" time used by AAPS.

Neighboring the AAPS\_logs is the Emulator file where the 5 downloaded files went:



- **10.1.3** Create a "start emulation" button on your desktop
- One of these files is "emulator\_GUI.py"
- Create a link to it
  - Drag that link onto your **desktop**
- Name it something like Emulator start.

- 10.1.4 Fetch two more .py files
- Repeat steps 10.1.1. and step 10.1.2. for 2 more py files from <a href="https://github.com/ga-zelle/Scan-APS-">https://github.com/ga-zelle/Scan-APS-</a>



Include these in your Emulator file (as in example shown in 10.1.2 above).

#### 10.2 Analyzing loop decisions in logfiles

89 Instead of making many screenshots every 5 minutes after meals, and analyzing them later, a much more

90 elegant and powerful way to analyze your loop decisions (and how you might want to influence them with

91 different settings, see <u>section 10.3</u> for this), is to use the Emulator.

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10.2.1 Set up a "no change" .vdf file.

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It is just empty in the lines that would define any change to be investigated.

Note: for "what-if" analysis, entries will be made (in a second .vdf later, see section 10.3)

To do this, just open **Notepad++** (from list of all programs on your PC). Store that in a file of your current

98 emulator project you are about to start (see my storage path in top line here)

99 The no change .vdf should look like something like this:



→ Dokumente → B\_Looping → Logfiles\_Emulator → Emulator\_Studien

autoISF\_factors\_ <- regarding this, see section 10.3.3.3 en

📋 🗐 🖻 🗓 🗘 Sortieren 🗸 🗏 Anzeigen 🗸 🚥

Studie\_1

Studie\_2

Studie\_3

noChange.vdf

Erase any entries after CR LF and also in lines 2 ff, if any

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Store this noChange.vdf in your Emul.-studies file (neighbors

 $104\,$   $\,$  your Emulator and Logfiles files). From that position, you

105 always make a copy and paste into each Study 1 ... n

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#### 10.2.2 Locate relevant logfiles

110 Make sure you have the AAPS logfiles that you want to analyze in an immediate-neighboring file to

111 your Emulator-- and to your Emulator Studies- files (File structure as suggested and shown

112 above).

113 Copy (not: move!) your noChange.vdf (see above) also into your Study file (must be in all of them).

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#### 115 10.2.3 Prepare the Emulator

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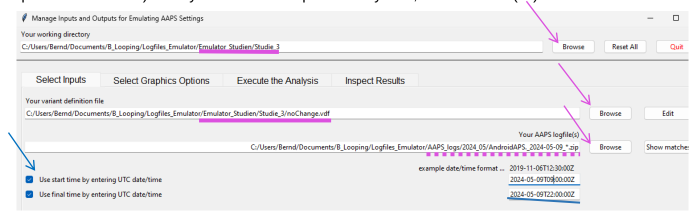
Now go to your PC desktop, and start the emulator by just pressing the button "Emulator

118 **start**" that you installed in step 10.1.3

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This opens a big dialogue box with 3 fields that you must fill in with the applicable path (without any

quotation marks "..") from your Windows Explorer file system, best done via (3x) Browse button:



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a) The top box marks the path to your current emulator project ("Studie\_3" is where I want to store results)

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- b) The middle box marks the path to your current vdf (what kind of analysis; here:
- "...noChange.vdf" = read-only; see section 10.3 for what-if)

c) The third box marks the path to your AAPS logfiles you wish to look into. A good way to do this is:

- Browse in your Windows Explorer to any logfile from the desired day (2024-05-09 in above example)
- Replace the time with an asterix \* (this means you look at all-day data, in UTZ time).
   Check whether this will work by pressing Show matches.
   You should see all logfiles from that day in a pop-up info box.
- As I wanted to look at 11 am –midnight (for lunch and dinner related data), I:
  - o clicked the bottom left two boxes
  - o copied the date 2024-05-09 over the default date in the bottom right two data fields
  - o after T (for time), I entered the desired time of analysis AFTER conversion into my local time (Central EU summer time minus 2 hours = UTZ; so to look at 11 to midnight of my AAPS screen, I must enter here 09.00:00Z, and below it 22:00:00Z).

Entries at the bottom are not mandatory, but when clicking these little boxes (bottom left) you can define a start and/or an end-point for analyzing, within the logfiles specified in the field above.

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146 10.2.4 Run emulation

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148 Now we are ready to go: Press "Run Emulation"

This produces sometimes an error message (e.g. if you have a syntax error, or incompatible software

versions: => seek help, in the Github materials provided by ga-zelle, or in Discord/Full-Closed-

Looping/emulate-aaps here: <a href="https://discord.gg/n3tD5eXExC">https://discord.gg/n3tD5eXExC</a>

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After a short moment results should show up, which you can look into in a couple of ways:

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## 10.2.4.1 SMB tab contents in (date..) **noChange.txt** result file

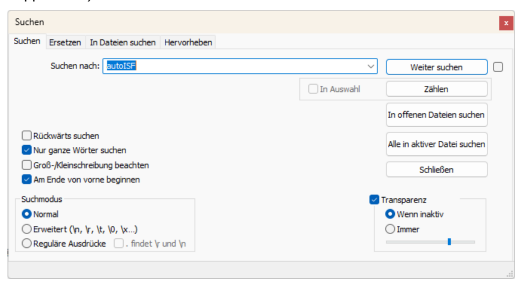
156 This basically gives you "all the SMB tabs" without needing to make screenshots every 5 minutes.

Search options help find what lines are of interest to your analysis:

By using the search function you can jump, in that super long list, to all places that e.g.

have "autoISF" in it or "script debug", or "SMB disabled" (if you want to analyze when that

happened...)



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10.2.4.2 Table of results (...noChange.csv file)

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The .csv file in your project folder gives a tabular presentation of how parameters like bg, iob,

iobTH, the various ISF contributors, bg target, insulinRequired etc. develop every 5 minutes, and

167 what SMB size and %TBR resulted.

168 It is a vast table, so you may want to reduce it to something more "digestable":

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170 10.2.4.2 Analysis of the (date..) **noChange.csv** table in Excel or LibreOffice calc.

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Best copy the entire table into a new .xls or .ods sheet, where you can:

add right next to the standard world time your corresponding "AAPS time"

<ul><li>174</li><li>175</li><li>176</li></ul>	For instance, adding +2/24 translates the UTZ column into central European summer time column next to it (where currently a row of Z stands). Likewise, subtract like -5/24 from UTZ for an US East Coast time scale.
177 178 179	Highlight all time fields (the entire columns), and switch from hh:mm:ss format to hh:mm.  (While the seconds are important for the loop's calculations, for our comparison with Nightscout or other charts and data, it is much easier without the seconds attached)
180	hide any column you find less important to look at for your intended analysis
181	That way, "boxes" (data fields) retain their original position in tables
182 183	Also, in case later you want to look into additional info, you can simply un-hide the relevant columns or time lines:
184	hide lines (time segments) you find less important to look at for your intended analysis
185	
186 187 188 189	Usually you will color mark where relevant SMBs were given, which of the ISFs (and underlying weights) was strongly contributing (note that this can be good or not good). Also where iobTH was exceeded, whether an Automation kicked in e.g. setting a TT, when there were periods with zero insulinRequired.
190 191 192	In <u>section 10.3.4</u> we present an extra tool that does a standardized table reduction and color marking for you!
193	You may be able to formulate a hypothesis or two, what settings (ISF_weights, iobTH%,
194	SMB_range_extention, autoISFmax) should be changed for improvement (then go to <u>10.3</u> )
195 196 197 198	10.2.4.4 Graph <b>noChange.pdf</b>
199	When you open the emulator, before executing any analysis, you can make settings under "Select Graphics
200	Options".
201	Select your preferred way of outputting decimals (point or comma).
202	Then select whether you want "All" possible outputs in the graph, or "Most" = all except those you tick "off"
203	in the boxes for each output parameter.
204	If you would use "Some/just a few", you would have to tick those few you that do want to see, by ticking the
205	corresponding boxes.

206 Recommendation is to look at (nearly) everything offered (as your default setting that you can leave 207 untouched in most of your emulator runs):

Manage Inputs and Out	tputs for Emulating AAPS	Settings			- 0
Your working directory					
				Browse	Reset All Quit
Select Inputs	Select Graphics	Options Execute the Ar	nalysis Inspect Resu	lte	
	· · · · · ·	Execute the Ai	inspect Nesu	ito	
The resulting graphics re					
All/-pred/-flowd	hart				
Select the decimal symb  ightharpoonup use ","  ightharpoonup use ","	ol for output tables	Coarse grained selection of grap     just a few     most (i.e. All but a few)     All  Fine grained selection of items to insulin chart content     Hide insulin required     Hide max bolus limit     Hide SMB     Hide basal rate	·	activity se ratio	Flowchart ON/OFF  Hide flowchart
			☐ Hide range p☐ Hide best slo☐ Hide other sl☐ Hide other sl☐ Hide best pa☐ Hide other p.	ope lopes rabola	

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After your emulation run, under Inspect Results, you can open the pdf file that is last in the results list offered.

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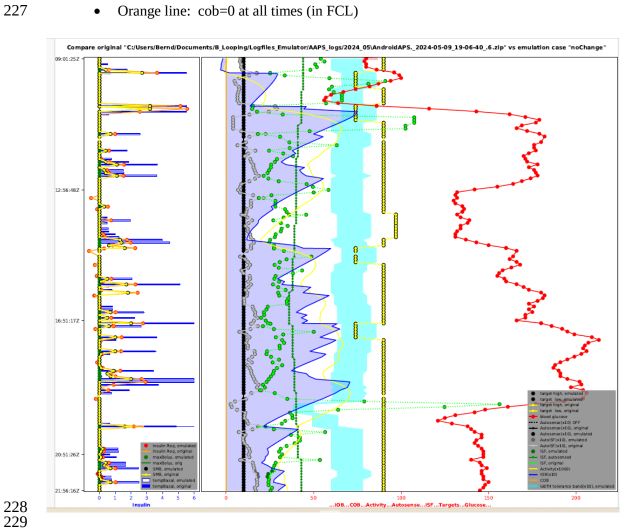
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- This **noChange.pdf** is a chart that shows along the time axis (down), from right to left:
- Red: the bg curve
- Yellow: the bg target (note that I do no manual "EatingSoonTT" but for bg rises over +10 mg/dl
   I have an Automation that sets low TT for a couple of minutes)
  - Light blue corridor: Left edge is set iobTH, and bandwidth +30% (would be +20% at elevated TT)
  - Dark blue line: iob (exceeding twice the iobTH, with temp. SMB shut-off
  - As bg did not convincingly come down enough, one could hypothesize that iobTH should be elevated. ((But, again, this would have to be confirmed also with other kinds of meals)).
- Thin yellow line: Insulin activity
- Green dotted line: ISF as would result from AAPS w/Autosens
- Green scatter points: autoISF ISF no Chage (lighter points) or what-if (darker points)
- Black line: Profile ISF

Gray scatter points: ISF weakened (to the left of black line) or strengthened (to the right)

Orange line: cob=0 at all times (in FCL)



More see discussed together with (yourChanges).pdf in section 10.3.3.4

237 238 239	"What-if" analysis using the emulator			
240	In the following you see an example how you can analyze a day of logfiles, and selecting the time span of			
241	interest, for instance 11-24 h to look at how autoISF managed lunch and dinner.			
242				
243	You have to go through the emulator twice. You already did it (section 10.2.) using the no change.vdf, and			
244	now start over with the same emulator with your (desired changes).vdf. Repeat, if you have two or more such			
245	vdf defined. (Just clear results before executing analysis each time. No worries, all results are automatically			
246	captured for all runs, all in your selected study file).			
247				
248	10.3.1 Define your investigated changes in a, or in several, (yourChanges).vdf			
249				
250	Define for which one to max three parameters in your current profile settings you want to look into a			
251	different setting. Recommendation is to use a factor, like for example current setting * 0.9 , or current			
252	setting * 1.2, and use that in your naming for this vdf file, too.			
253				
254	Within the same study, you can make several runs with several vdf files.			
255	All results, like the csv results table, will appear then several times in your study file, only with different			
256	name endings as in the underlying vdf.			
257				
258	Example: I like to check in my actual data (they are in my noChange.vdf emulator run <b>), in which time</b>			
259	<b>points the following parameter changes would make a</b> (how) big <b>difference</b> in the loop's decision:			
260	• 20% higher bgAccel_ISF_weight to boost the first SMBs stronger: How would that tend to ramp			
261	up early iob; and might that get too strong in other parts of the data? Or does it bounce into a			
262	restriction (maxSMB size; autoISFmax; iobTH) that I might need to widen?			
263	<ul> <li>Doubling my cautiously set bgBrake_ISF_weight shall give me insight into the workings of that</li> </ul>			
264	parameter (and whether using a much smaller weight than for bgAccel_ISF_weight is really			
265	what I should keep doing)			
266	• As my bg came down from a persistent high quite slowly, I elevate the dura_ISF by 20%			
267				
268	Actually, it would make more sense to first find my "optimal", maybe indeed elevated,			
269	bgAccel_ISF_weight. <i>Then</i> , do a noChange (!) run <b>with that,</b> plus a (yourChanges) run with the stronger			
270	dura weight, investigated on that basis.			
271	Reason: 1) As we always say, better do only one change at a time. 2) A better job with bg control via			
272	bgAccel_ISF will reduce the peak height and provide a different (easier) scenario for dura_ISF to manage.			
273				

Now, to **write** your **(yourChanges). vdf for the emulator** (this is same procedure as you did in section 10.2.1 for the noChange.vdf):

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- just open Notepad++ (from list of all programs on your PC) to create a new vdf.
- Alternatively you can also take another pre-existing vdf file, and copy it into your current project giving it a new name (re-name it)
- name your vdf (in our example: 1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.vdf) ...
  - ... and store that in a file of your current emulator project you are about to start (see my storage path in top line here)

**Caution:** Make absolutely sure (best by looking it up in the SMB tab, down in the profile set section) to **spell each term exactly** as your loop uses it (probably w/ decimal points, not comma)

• ...when you make one line per parameter (separating entries with spacers->):

profile->(parameter) ->->profile['(parameter)']\*(factor)->->###(comment as you like)

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The (yourChanges) .vdf should look like something like this:

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```
1.2_bgAccel_2.0_bgBrake_1.2_dura.vdf AndroidAPS._2024-05-09_02_00-31_0.zip.orig.bt 2 = 2024-05-09T09-00-00Z.1.2_bgAccel_2.0_bgBrake_1.2_dura.txt 2 = 2024-05-09T09-00-00Z.1.2_bgAccel_2.0_
```

CR = LF= Erase any entries after CR LF and also any entries in lines below, if any

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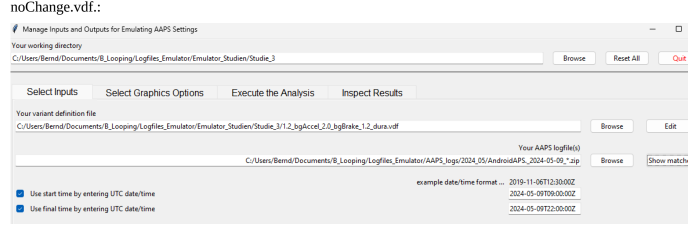
10.3.2 Run the emulator with (yourChanges).vdf

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The "what-if" emulator run is done the same way as you did the noChanges run (<u>section 10.2</u>), however, now, the **(yourChanges).vdf** must be loaded into the 2<sup>nd</sup> input field, where formerly you had the



- 301 In the 3<sup>rd</sup> input field, give the path to your stored logfiles. A good way to do this is:
  - Browse in your Windows Explorer to any logfile from the desired day (2024-05-09 in above example)
    - Replace the time with an asterix \* (this means you look at all-day data, in UTZ time). Check
      whether this will work by pressing Show matches. You should see all logfiles from that day in a
      pop-up info box.
    - As I wanted to look at 11 am –midnight for lunch and dinner related data, I:
      - o clicked the bottom left two boxes
      - o copied the date 2024-05-09 over the default date in the bottom right two data fields
      - o after T (for time), I entered the desired time of analysis AFTER conversion into my local time (Central EU summer time minus 2 hours = UTZ; so to look at 11 to midnight of my AAPS screen, I must enter here 09.00:00Z, and below it 22:00:00Z).
  - After making these entries, press Execute the Analysis, (evtl also Clear old Data) and then press Run Emulation, I can look the results up under "Inspect Results":

#### 10.3.3 Emulation results

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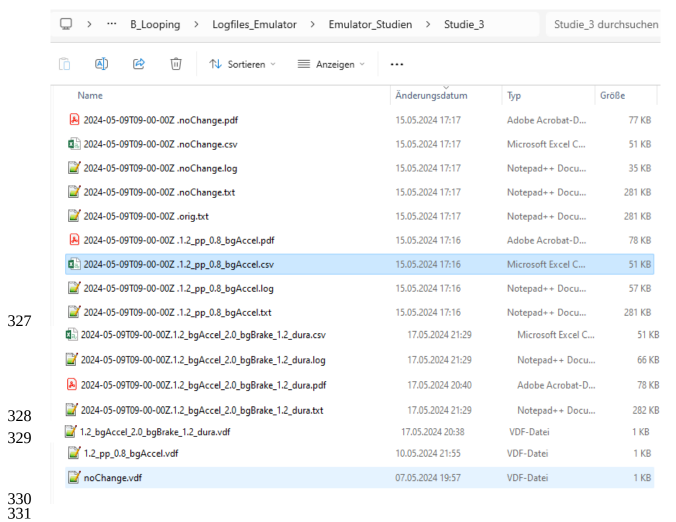
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324

325 326 Your working directory C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie\_3 Inspect Results Select Inputs Select Graphics Options Execute the Analysis \*.log - Your file showing edits from the variant assignments  $C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie_3/2024-05-09T09-00-00Z.1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.log$ \*.csv - Your table comparing key values of original vs emulation C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie\_3/2024-05-09T09-00-00Z.1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.csv \*.delta - Your table comparing bg deltas of original vs emulation  $C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie_3/2024-05-09T09-00-00Z.1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.delta.pu$ \*.orig.txt - Your short log of original analysis C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie\_3/2024-05-09T09-00-00Z.orig.txt \*.txt - Your short log of emulated analysis  $C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie\_3/2024-05-09T09-00-00Z.1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.txt$ \*.pdf etc. - Your graphic file comparing key values of original vs emulation Browse C:/Users/Bernd/Documents/B\_Looping/Logfiles\_Emulator/Emulator\_Studien/Studie\_3/2024-05-09T09-00-00Z.1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.pdf Show

All results from your (yourChanges).vdf emulator go automatically where the noChange.vdf results are already stored, in our example into the "Studie 3" file, below:

Besides the 1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura.vdf case which I like to look into for the present high carb meal, I also prepared another vdf that investigates a factor 1.2 stronger pp\_ISF and a weaker, factor 0.8, bgAccel\_ISF (with the intention to test this, and a noChange, on a low carb meal later.



10.3.3.1 Logs check: log vs txt

The **noChange.log** has all the info your series of SMB tabs had that day.

How to search in this vast list is shown elsewhere (see section 10.2.4.3).

Likewise, the **(yourChanges).log** gives for each loop decision in all detail how and why that decision would have changed with the different parameter inputs you are checking out here

In the two examples here, , it was a check on the difference

- a 20% stronger pp\_weight and 20% weaker bgAccel\_weight
- a 20% stronger weight for both, bgAccel\_ and dura\_ISF, and a doubling of bgBrake\_weight

342 would make.

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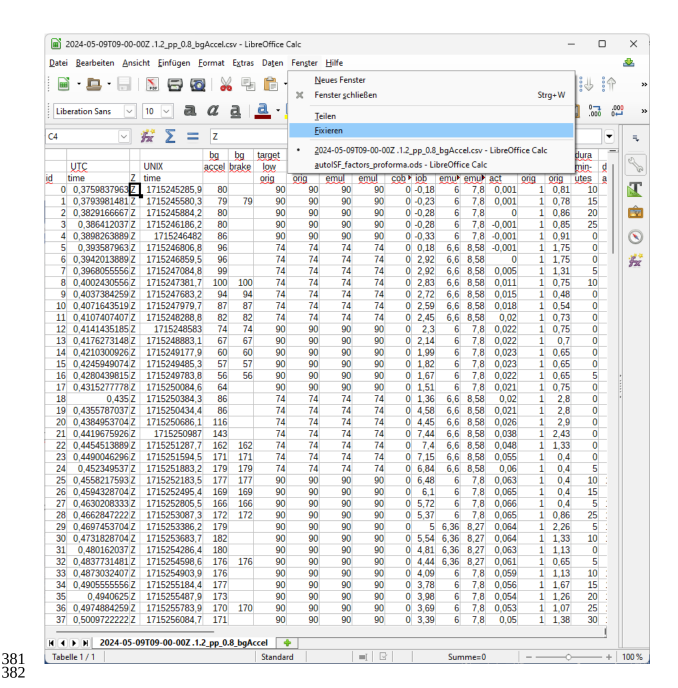
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Note that all these "what if" data can only give rough hints, notably about **the first** greater change that you would see with the investigated changed setting. So it works well for investigating how to ramp up iob quickly after detection of acceleration.

However, any relevant change would put your bg curve on a different trajectory, so that would influence all following results. Therefore, what you get here is **not** a complete modelling how your bg would have developed in the alternative scenario.

350	But you can investigate in which stages the parameter(s) you are looking at in your current "what-if" had big
351	influence, and in which direction the changes would go. (see also charts shown in section 10.3.3.4).
352	Analyzing how to safely come down from a high glucose plateau while limiting hypo danger towards the end
353	of digestion is also to some extent possible.
354	
355	A good other way to employ the what-if analysis is real time, on your smartphone, using speech synthesis
356	(see section 11): Then you get real-time info, as to exactly when a significantly different proposal would
357	emerge, and can decide (and watch!) real-time whether to follow the new idea and not was probably better.
358	
359	Observe that a setting change must work well for you
360	<ul> <li>not just in one point of time, and</li> </ul>
361	<ul> <li>not just for one kind of meal,</li> </ul>
362	but you must look at all time slots in the investigated meal, plus analyze with the same tool a totally different
363	meal within your usual spectrum, how things work out there
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365	10.3.3.2 Tabular results
366	
367	The <b>noChange.csv</b> table gives all relevant data. Besides development of bg and iob you see the calculated
368	insulinRequired in each loop decision, and how each of the autoISF categories contributed to the decision
369	(notably regarding SMB size).
370	
371	The <b>(your changes).csv</b> shows in detail how <b>every single</b> loop <b>decision</b> would be influenced by the different
372	settings you are investigating. To inspect that huge table, click on the Z behind the start UTC time entry
373	(see black box in the Z column of the following table
374	If you like to see the bg in each screen, too, go 3 or 4 columns farther to the right with your black
375	box.
376	Then, go to window/fix. Now you can scroll through the data and always see headline and time (or time and
377	bg level).
378	To further ease analysis, feel free to temporarily erase (hide) any columns that you (think you) do not
379	need for the intended analysis. More suggestions see in section 10.2.4.2
380	



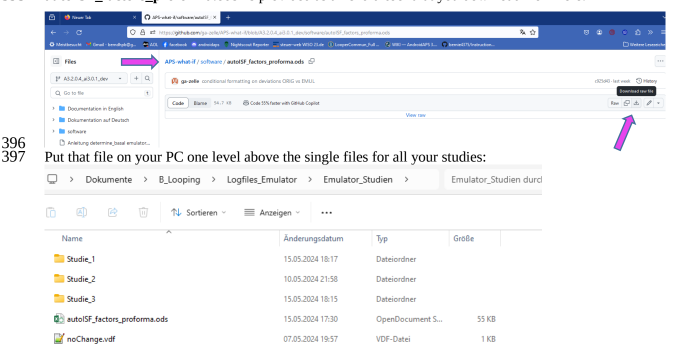
Still, the csv tables are overwhelming. You could proceed in either of two directions now:

A) Convert both (or all 3) csv files into one table in Excel or into Libre office calculator. Hide columns (and eventually also lines) that are of no particular interest for your analysis. Mark differences between noChange and (yourChanges) column data with color, add extra columns with additional calculations ...

This route is good to compare quantitative impacts of autoISF categories in critical time points.

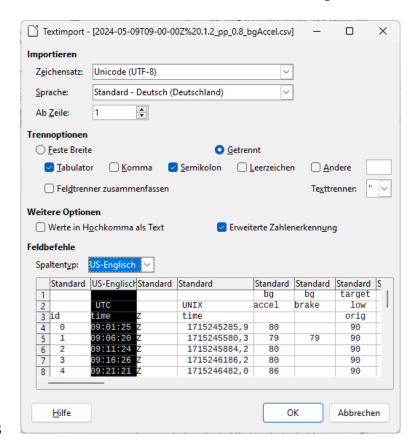
B) For the core data relevant to assessing your autoISF settings, there is an extra tool for convenient analysis (see next section)

## **autoISF\_factors\_proforma.ods** is provided as an **extra tool** that you download from here:



Now, if we want to use this tool on the two csv files of our Studie\_3 file, we must proceed as follows (for *each* of the two .csv files, *separately*):

- 1. Click on the .csv file and open in Libre office calculator.
- 402 2. Make sure the time column is set to US\_English:



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3. Now start, in Libre office calculator, the autoISF\_factors\_proforma.ods ...

This turns the first 30-some lines of your csv table (left side) into a form in which important effects are

highlighted in color, and formatting is improved:



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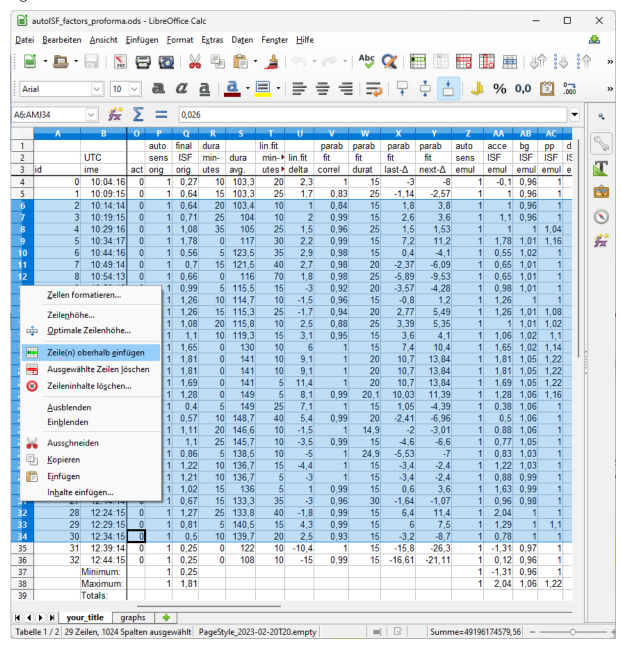
Now, you want this for the entire table.

In the autoISF\_factors\_proforma table, highlight 20 or more lines (not including the first or last), and mouse

412 right hand/insert above ...

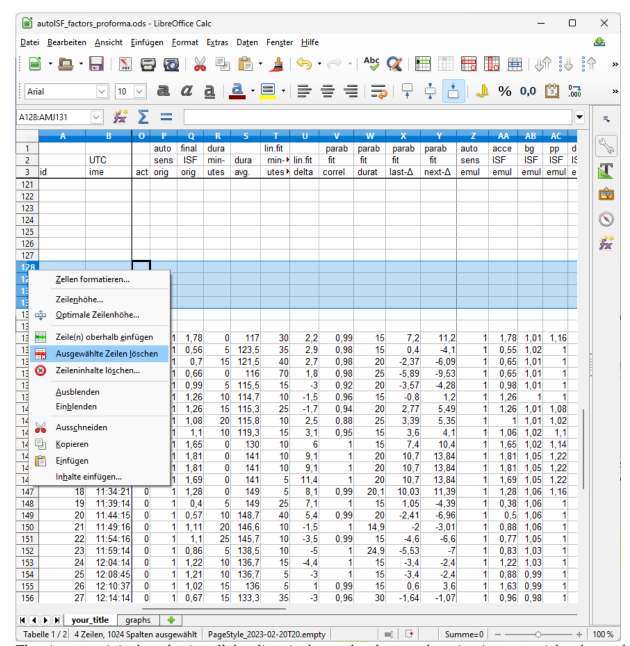
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Do this as often as you need to create the number of lines that your emulated csv file comes with.

416 If you ended up with too many lines, erase the superfluous number (any four, in the example):



Then just copy it in, by selecting all data lines in the emulated csv, and pasting (paste special, values only)

into box A4 of your "elonged" autoISF\_factors\_performa.ods.

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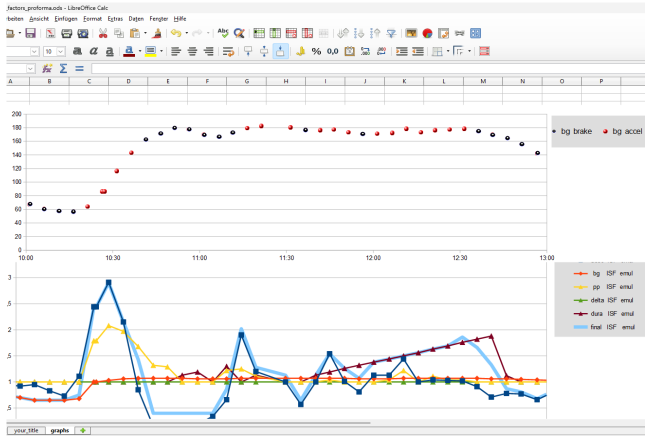
The bottom tab "your\_title" should be re-named by you, best with day of log you analyze, and your what-if parameters (so, the name of your csv file could be put in here)

Now you have a table with optimized lay-out that incorporates key data from both your no change AND of your investigated changes.csv files.

A super neat extra feature is already pre-programmed, which you can see if you click on the bottom **tab** "graphs".

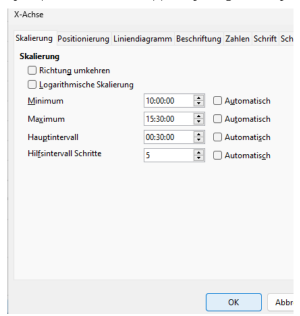
428 The top graph is the bg curve (the actually seen bg).

Note that for the what-if no bg development over the time range is available. (The noChange one is also given there).



The bottom graph (do one for each, the noChange or the (yourChanges) case) shows the amplification factors coming from each autoISF category, and the overall resulting ISF amplification.

You probably have to widen the time scale (double click on the time axis, and type the desired time span (min and max UTC)(and spacing of data points, 00:30:00 or 00:15:00) into this box:



In the given example above, the 2.5 hours were not enough yet to analyze this 10:30 UTC (12:30 AAPS) lunch; we need to look until bg is near target (hopefully before dinner starts).

A similar graph is available on the (i-)phone if you use the autoISF dev variant of iAPS (and maybe of Trio, in the near future). (Later insert details here, or in section 11)

442 443	10.3.3.4 Chart coming with the Emulator
	In case you find the cuture stone described in the proceeding costion "too much" place the appulator offers are
444	In case you find the extra steps described in the preceding section "too much", also the emulator offers one
<ul><li>445</li><li>446</li></ul>	chart (the pdf offered at the bottom of the screen as shown below the "10.3.3 Emulaton results" headline).
447	First look at the initial bg rise in the noChange.pdf chart (emulation results from your noChange.vdf run),
448	and see how bgAccel_ISF and pp_ISF acted, or could have acted in improved ways.
449	Then look into in (yourChange).pdf to see potential effects (or what other change to try). (Actually, you
450	probably will have to go into a detrailed analysis of several lines and columns of the tables as discussed in
451	sections <u>10.3.3.2</u> and <u>10.3.3.3</u> ).
452	
453	Note that ideally we want FCL coverage of our entire "normal day" meal spectrum by one set of
454	settings. So, not much is gained if you put a lot of effort in optimizing FCL settings for one
455	meal.
456	
457	You will need iterations. Do such analysis for two or three very different meals that you wish the
458	algorithm to automatically handle. See $\underline{\text{section 4.2}}/\underline{4.3}$ on how meals with very different carb loads
459	might benefit or also suffer from too aggressive or to mild (category)_ISF_weights you could set.
460	
461	
462	The initial iob received might be limited by allowed SMB sizes, autoISFmax, or the (dynamic!) iobTH. You
463	will have to look into the data table to find out about this (a quick orientation - notably regarding the light
464	blue iobTH band, see next page - is also possible in the pdf result files you have in your project file (project
465	file example "Studie 3" in $2^{nd}$ chart under the $\underline{10.3.3}$ . headline).
466	
467	Only once you found OK weights for bgAccel- and pp_ISF_weights, does it make sense to go tune the
468	$dura\_ISF\_weight.\ 12:00-12:45\ UTC\ in\ above\ graph,\ the\ resulting\ effective\ ISF\ is\ dominated\ by\ dura\_ISF.$
469	Just judging from the picture, a stronger weight might be worth trying. However, we really need to see the
470	insulinRequired calculation and the further development because impatience about bringing bg values down
471	faster too often results in hypoglycemia later.
472	
473	The <b>noChange.pdf</b> is a chart that shows along the time axis (down), from right to left:
474	• Red: the bg curve
475	• Yellow: the bg target (note that I do no manual "EatingSoonTT" but for bg rises over +10 mg/dl
476	I have an Automation that sets low TT for a couple of minutes)
477	• Light blue corridor: Left edge is set iobTH, and bandwidth +30% (would be +20% at elevated
478	TT)

Dark blue line: iob (exceeding twice the iobTH, with temp. SMB shut-off

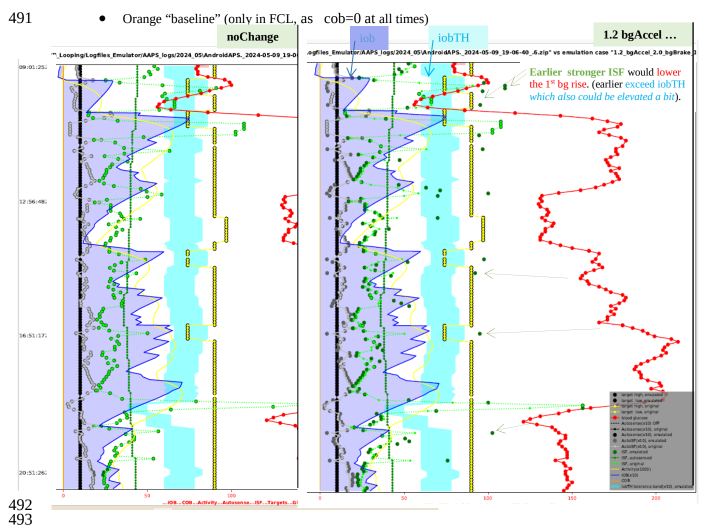
As bg did not convincingly come down enough, one could hypothesize that iobTH should be elevated. ((But, again, this would have to be confirmed also with other kinds of meals)).

• Thin yellow line: Insulin activity

- Green dotted line: ISF as would result from AAPS w/Autosens
- Green scatter points: autoISF ISF no Change (lighter points) or what-if (darker points)

Foreseeably, this is the strongest difference between our noChange (left) and 120% bgAcel\_ISF\_weight (right) in the picture below. (Note the red bg curve is *both times* the really seen bg, because the what-if case only looks at each single loop decision). The first ( -> ) time the dark green dot is far to the right, this *would* get the bg down, we *would start to see* a ( <- ) bg lowering effect, shifting the red curve to the left

- Black line: Profile ISF
- Gray scatter points: ISF weakened (to the left of black line) or strengthened (to the right)



Regarding the other changed parameters: Stronger dura\_ISF would suggest more insulin towards the end of plateaus; this should have helped in the 1<sup>st</sup> plateau (red curve, top right quadrant of the picture). However,

same setting would have to work also on 2<sup>nd</sup> plateau; the chart cuts off there, so too early to see whether a hypo danger might result.

Effect from doubling the bgBrake\_ISF effect are hard to evaluate. Better probably to look in .csv tables, or run a separate emulation for that change only.

Always check for 2 or 3 kinds of your meals whether the "new" parameter settings really are on average better. (See negative example in <u>case study 8.2!</u>)

Part of both above shown charts (left side of each, with blue peaks) was cut out.....

(Unfinished / to be explained later) (...note: yourChanges = 1.2\_bgAccel\_2.0\_bgBrake\_1.2\_dura)

