Metabotools tutorial II - Integration of quantitative metabolomic of

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INTRODUCTION

In this storial, we generate a controlucitized model by integrating quantitative extraoelular metabolonic data (1). We will alterwards analyze the behavior of the models to

satisfaces in this by using phenotypic phase plane analysis.

Before running a section in the babrial, read the corresponding sections in the Ministo Yooks protocol and supplemental babrial (bata wheet 3;

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PROCEDURE
Clear workspace and initialize the COSPA Toolbox

clear

ctear Inittabrañas bas

Step 0 - Define the output location and set the LP solver

saleer = 'gardo'; % can be quroi; or 'ibn saleer's 'changeCobrataleer(colver, 'IP'); If columnity = 1

display('The LP colour is set.'); else erwr('The LP colour is not set.')

Load and check that the input model is correctly loaded

teterialment = filegarts(which("teterial_menactoster.e")); if exist([teterialment filecop "ctaring_model.ast"], "file") == 2 % 2 meas it's a file.
ctaring_model = meastmadel([teterialment filecop "ctaring_model.ast"]);

errar('The ''ctarting_model'' could not be loaded.');

Check output path and writing permission

if =exist(outputPsth) == 7

to Make and case dummy file to test writing to output directory

A = rand(1);

try care([outputPath fibecep 'A']); catch ME

end

Step 1 - Shaping the model's environment using setMediumConstraints

Define the model bounds using setfled/unrConstraints
Constain the model abounds using the state setted to readure corporation. Yorks and, define the set of exchange reactions for which examendations data are available. In this examend, not find any available.

example, no data are available medium_composition = {}; met Canc aM = {};

Define containts or basic medium components (i.e., metabolins that are uprate from the medium but noncaptured by the measured distay and distay

last = ("EC_CAT(o)", "EC_Cb(o)", "EC_Cb) mediumCampounds = [lost mediumCampounds] mediumCampounds_lb = -180;

Define also additional commands to that the model technique p_{ij} , according of again, essential amino additionate onto the taken p_{ij} destinational commands $= \{ (B_{ij}(x_i)(x_i)^*, (B_{ij$

"REJECTIO", "REJECTION", "REJECT

Apply the medium constraints previously defined using welderSurvivorsations. Note that you can also provide internation initiated to the cell concentration of insight (politicity); the Street js, the current value and the new value for infinite constituents (perpectively current, inf and set, infi.

or INSIGHT S. INSI

cellmeight = []; t = []; set_inf = 2000; current_inf = 1000; close_exchanges = 0 [notellmeight | notell

[modelPedium, basicMedium] = cetMediumicnotraints(ctarting_model, cat_inf, current_inf, medium_composition, mat_ctar_inf, cat_inf, current_inf, medium_composition, mat_ctar_inf, cat_information_inf, mediumicnomedium_composition_information_composition_information_composition_information_composition_information_composition_information_composition_information_composition_information_composition_compos

Step 2 : Generate an individual exchange profiles for each sample

Generate included sprace and excending purplies from fluxes data using prepiritergozionicidant Note that negative flux values are interpreted as uptake and positive values are excending. Moreover, the function removes from each sample the more distribute updates or excending the cannot be remobile by the resolved due to missing production or engagedation patterns, or blooked resolved, and because it only excending to opposition, only excending is definited from the earlier printing enhancing updates and excending the desired of the engaged.

laad([tatarial@sth filecep 'tatarial_ff_data.est']);
sode(= sode(bediam)
text_pia = 500;
text_pia = 0.0001)

prepintegrationQuant(model, metiata, exchanges, samples, test_max, test_min, output Charmars -EXCEPT modelMedian samples tol saleer outputPath internalPath colverQuan

Use checklicitangeProfiles generate a summary of the number of uptake and secretion exchanges per samples.

ments = mul | Rapped_cothanges, mints, mapped_uptaw, mapped_cocretion] = checksushangerutites(samples, outputPath, maets); | Chargars = EXERT modelfedies samples tal saleer mapped_cochanges_outputPath tatorialPath objectpoint

Step 3 : Generate contextualized model

Use the function certificated containing is integrate the update and excention profites and generate condition-specific mentation models for each sample. The function allows the self-look of an efficiency of the function allows the self-look of an efficiency of the function allows the self-look of an efficiency of the function of th

changetabrabolowr(saleerQuart, "IP");
mindrawth = 0.000/n lower bound to the biomacs reaction obj

ab) = 'bloade_maction3';
as_servine = ('BX_SD(e)');
as_spine = ('BX_SD(e)', 'BX_SDAZ(e)');

magname to tactors; all SISIO()))
medium = ();h reactions that should be excluded from mini
tal = 10-6;
model = modelMedium:

AdMictished = ("Elliscolia"), "Elliscolia") by metabolic exchange that are added to the upper and lawer admictished radius = 13p flow values that are added to the upper and lawer bound; [Resolitatified] laws, deverowanted() = organization(sizition(sai), saiples, to), andersoft, sby, on_excertise, ...

No. prise, medium, additionals, additionals,

[Re_abset_all_water] = caristicondomenoaque(Recultollites, campled) clavare-futtor moltonia camples resultationis devicements to passe_all_unique tal salver appet_enthages output/973 t

CAMP[CAMPATH Tileop "CANADA"]);
classes—CROT and Delias capts neutralizations deviaments til siles appel extages outpoten totalization
(Dep 5: Analyse has to descending once

Use the function analyzedingle/dene/Celetion to predict and analyze the sents of essential genes across a set of models.

ignore, your Continuitions, portrainments) - analyses applies applies and results and institutes, unique to the property of th

Step 6 : Check reaction essentiality
Use the function checkfiles/Puriti to investigate which individual gene-associated reaction makes the model inhealths (i.e., macrions associated with a gene need to

genec_ta_test = ("NITELL'); viet of genec
[PER_Mans_NZ, ListResits] = checkeffetExcetD(samples_ta_test, fill, genec_ta_test, samples, mesuitsEllLines);

Clearant - CECTY maintenance usuplex results tabilities cover/develope by about 11 major genes PRU, Nort, NO Estimation to Step 7: Generate an intersect model and an uniform found to the common section and common section model common model (normalized and an intersect model (downstandard) and an intersect model (downstandard). Additionally, it will be set of reactions

of Tables in adultations produces (offsekling that distinguist the union and the research model for further analysis.

Million at 1)

Million (1)

M

samples to test a sampless wood-burt of models

model = starting_model;

sive((output/eth filecep 'cuminy'));

Step 8 : Predict differences in metabolite production or consumption

Use the function prescribulaçiate is upwast the production or consumption plateal using all of mentiollates of invest (motified a 4,0) in which make defined by an application of the control production of the control production and the prescribe motion (specified which the control production that the control production control is mentioned by an application and the production and the production and the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control is set of the production control in the production control

numbolite of interver (maximum combusing on, maximum combusing flar). This analysis requires a quadratic programming (QP) solver.

changes duration verifications, "(2") is
all to "(2) a color (1).

camples = samples(1:4, 1); dir = 1;

Example A - ATP production

% exclude transport reactions from flax split analysis
transportman = ('ATPG') 'ATPG') 'ATPG'; 'ATPGG'; 'ATPGG

[mult, resittationthines, sectorit, sasissa_contributing_ras, sasissa_contributing_Two, Attyrial] = predictFlactplits(sass), adj, sectiont, casples, resultabilited lines, dir, transportment, Attyrial, carban_contribution_co

clear ATP/rod transportRoss netZtect maximum_contributing_ron

Example B - NADH production

ampie is - MALIN production 121est = ("nam(s)", "nam(s)", "nam(s)", "nam(s)", "nam(s)",

[molt, mentanticultains, semmed, memoricultains, ros, desime_contributing_flow_man() = predictFlowlylits(made),...
mby_metrest_man() = mentanticultains, dir_fraequetman()
max_p memoricultains_man(, m);
max_p moderners_man(, m);
max_p

Example C - FADH2 production

transportance = ("reprotor" | "reprotor");

transportEase = {"MONOTIV"; "MONOTIV"}; wtZtest = {"famis(c|", "famis(n|", "famis(n|", "famis(n|", "famis(n')");

[Well, Resittationities, second, assism_contributing_rm, assism_contributing_flow_PARMO] = predictFloodplits(socie_... alp, settlect, caples, Resittationities, dir, transportance) PM = [PM section_Contributing_contr, 1]];

Example D - NADPH production

transporttess = ("stometra") | "stometra");

wiffest = ('mage(c)', 'mage(e)', 'mage(e)', 'mage(c)', 'mage(c)

[mult, resittatifoltines, sectuals, assuma_contributing_res, maxima_contributing_five_MARTM] = predict(lactplite(model,...de), sections, complex, resistatifoltines, dir, transportment)
THE = [PM maxima_contributing_resi(, 1)];

save([output#ath fileses 'flux[slitt']);

Step 9 - Blustrate the phenotypes (PHs) on 3Dplot
The function naived/size allows illustration of the results of the armious analysis. The calons specify different open open.

make Explot (PMC, maximum_contributing_flux_ATP, forts, output FWTh, diff_view); Step 10 - Perform Phase Plane Analysis

Use the function perform PPP to investigate the behavior of the models to variations in flux through a pair of exchange reactions. The billowing example illustrates the values to be treated is defined by the number of steps and step size (step num and step size). The direction of exchange is defined individually for each exchange (direct) $\mathbf{mets} = \{ (\mathsf{max}_{i}) | (e) \land_{i} \land (\mathsf{max}_{i}) \land_{i} \land_{$

[Recultablicalizates] = performPPP(Recultablicalizates, mets, steg_size, samples, steg_num, direct); save([outputFath filecep "FFF"]); Use illustrate ago to illustrate the results of the shape place analysis.

label = ("blucose uptake (feol/cell/hr)"; "Buygen uptake (feol/cell/hr)"; "browth rate (hr-s)");

camples = ('IDMEVE');
illustrate_ppp(RecultsAllCellLines, mets, outputFath, samples, label, fants, tol);

Science 336: 1000-1046.

REFERENCE 1. July M. Nilsson R. Sharma S. Madhusudhan N. Kitami T. et al. (2012) Metabolite Profiling Identifies a Key Role for Grucine in Rapid Cancer Cell Profilesion.