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function example_use
clear;clc;close all;

%initialize rates
beta = 1.1;
gamma = beta;
kinit= 2.1;
bs=10;

%initialize simulation parameters
nCells = 1e4;
rng(floor(log(4815162342)));
S = [bs 0; -1 1; 0 -1];
nT = 4;
kpar = [kinit,beta,gamma];
Tmax=5/min(kpar);
tvec = linspace(0,Tmax,nT);
t_matrix = repmat(tvec,nCells,1);

%run Gillespie simulation
X = gg_200110_gillespie_geom_1(kpar,t_matrix,S,nCells);

%initialize approximation parameters
N_approx_taylor = 8;
N_approx_laurent = 5;

dim = nT-1;
nrow=3;

tvec(end) = inf;

%visualize the Gillespie simulation results
for i = 1:(nT-1)
    %visualize mature marginal
    figure(1)
    subplot(1,dim,i)
    histogram(X(:,i
+1,2), 'BinMethod', 'integers', 'Normalization', 'pdf', ...
    'FaceColor', 0.5*[1 1 1], 'EdgeColor', 'none'); hold on;
    title(sprintf('t = %.3f', tvec(i+1)), 'FontWeight', 'Normal');
    xlabel('mRNA copy number');
    ylabel('Probability');

    %visualize nascent marginal
    figure(2)
    subplot(1,dim,i)
    histogram(X(:,i
+1,1), 'BinMethod', 'integers', 'Normalization', 'pdf', ...
    'FaceColor', 0.5*[1 1 1], 'EdgeColor', 'none'); hold on;
    title(sprintf('t = %.3f', tvec(i+1)), 'FontWeight', 'Normal');
    xlabel('pre-mRNA copy number');
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        ylabel('Probability');

        %visualize joint distribution
        figure(3)
        subplot(nrow,dim,i)
        histogram2(X(:,i+1,1),X(:,i
+1,2), 'BinMethod', 'integers', 'Normalization', 'pdf', ...

        'DisplayStyle', 'tile', 'ShowEmptyBins', 'on', 'edgecolor', 'none');
        hold on;
        title(sprintf('t = %.3f', tvec(i+1)), 'FontWeight', 'Normal');
        if i==1
            ylabel('Gillespie', 'FontWeight', 'bold');
        end
        if i==3
            yyaxis right
            set(gca, 'ytick', [])
            ylabel('mRNA');
            set(gca, 'ytick', [], 'ycolor', 'k');
        end
        xlim([0, max(X(:,i+1,1))]);
        ylim([0, max(X(:,i+1,2))]);
    end

    tvec(end)=inf;

    %compute and visualize numerical integral results
    for i = 1:(nT-1)
        M = max(X(:,i+1,1))+15;
        N = max(X(:,i+1,2))+15;

        %compute mature marginal
        Pa_marg = gg_200228_numint_geom_tdep_3(kinit, bs, gamma, M, N, tvec(i
+1), 'mature');

        %visualize mature marginal
        figure(1)
        subplot(1,dim,i)
        plot(0:(N-1), Pa_marg, 'b-', 'LineWidth', 2);

        %compute nascent marginal
        Pa_marg = gg_200228_numint_geom_tdep_3(kinit, bs, gamma, M, N, tvec(i
+1), 'nascent');

        %visualize nascent marginal
        figure(2)
        subplot(1,dim,i)
        plot(0:(M-1), Pa_marg, 'b-', 'LineWidth', 2);

        %compute joint distribution
        Pa = gg_200228_numint_geom_tdep_3(kinit, bs, gamma, M, N, tvec(i
+1), false);

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    %visualize joint distribution
    figure(3)
    subplot(nrow,dim,i+dim);
    [x,y]=ndgrid((0:(M-1))-0.5,(0:(N-1))-0.5);
    h= pcolor(x,y,Pa);
    set(h, 'EdgeColor', 'none');
    xlim([0,max(X(:,i+1,1))]);
    ylim([0,max(X(:,i+1,2))]);
    if i==1
        ylabel('Numerical','FontWeight','bold');
    end
    if i==3
        yyaxis right
        set(gca,'ytick',[])
        ylabel('mRNA');
        set(gca,'ytick',[],'ycolor','k');
    end
end

%compute and visualize analytical integral results
for i = 1:(nT-1)
    M = max(X(:,i+1,1))+15;
    N = max(X(:,i+1,2))+15;

    %compute mature marginal
    Pa_marg =
    gg_200325_analyt_geom_tdep_vec_31(kinit,bs,gamma,M,N,...
        tvec(i+1),'mature',N_approx_taylor,N_approx_laurent);

    %visualize mature marginal
    figure(1)
    subplot(1,dim,i)
    plot(0:(N-1),Pa_marg,'r--','LineWidth',2);
    ylim([min(ylim)/5,max(ylim)]);

    %compute nascent marginal
    Pa_marg =
    gg_200325_analyt_geom_tdep_vec_31(kinit,bs,gamma,M,N,...
        tvec(i+1),'nascent',N_approx_taylor,N_approx_laurent);

    %visualize nascent marginal
    figure(2)
    subplot(1,dim,i)
    plot(0:(M-1),Pa_marg,'r--','LineWidth',2);
    ylim([min(ylim)/5,max(ylim)]);

    %compute joint distribution
    Pa = gg_200325_analyt_geom_tdep_vec_31(kinit,bs,gamma,M,N,...
        tvec(i+1),false,N_approx_taylor,N_approx_laurent);

    figure(3)
    subplot(nrow,dim,i+2*dim);

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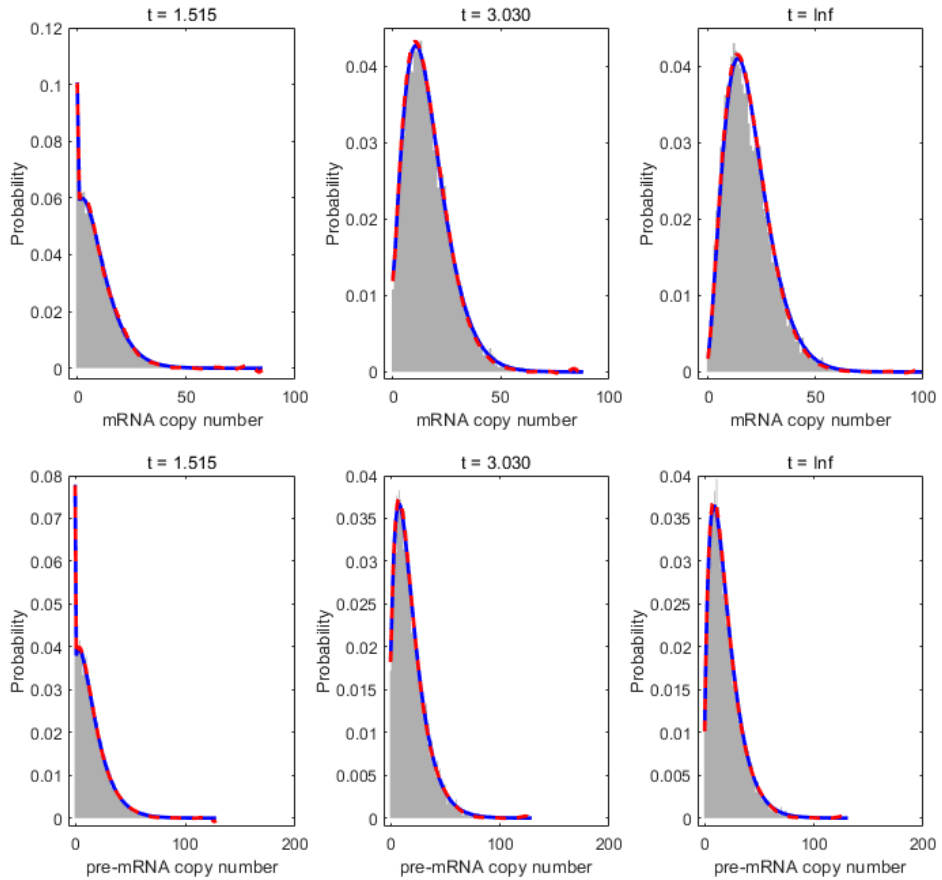
```

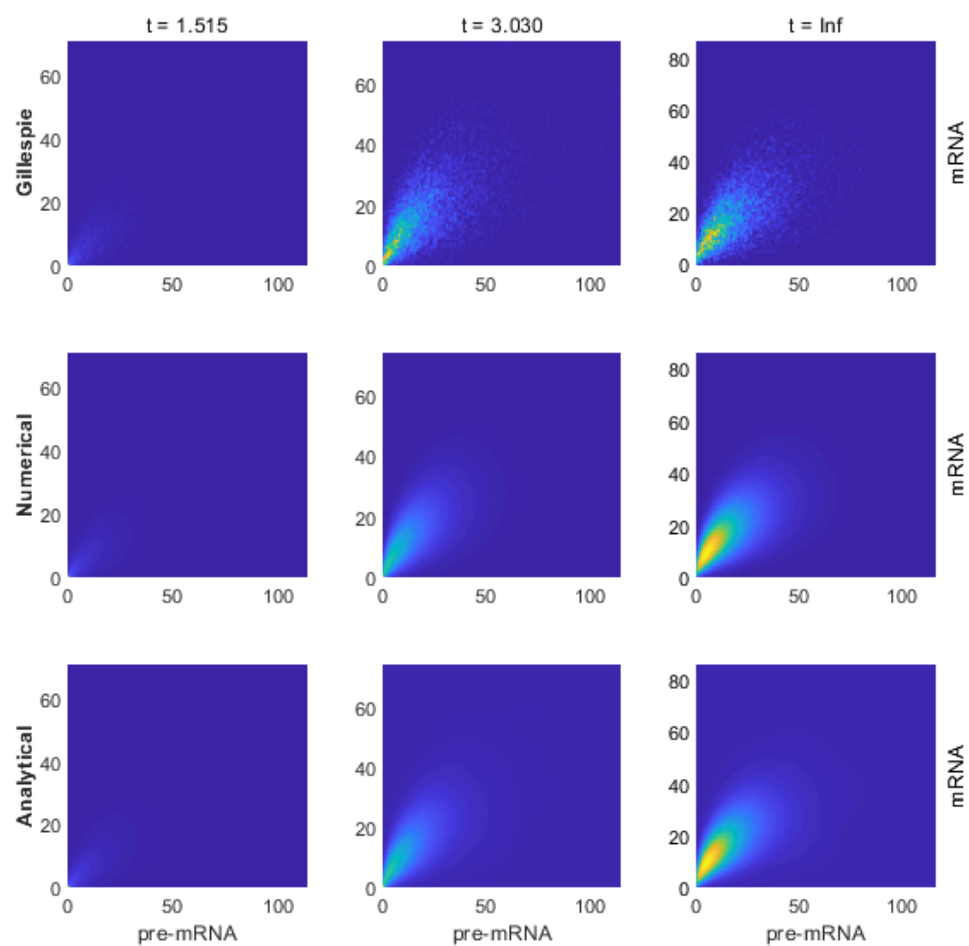
[x,y]=ndgrid((0:(M-1))-0.5,(0:(N-1))-0.5);
h= pcolor(x,y,Pa);
set(h, 'EdgeColor', 'none');
xlim([0,max(X(:,i+1,1))]);
ylim([0,max(X(:,i+1,2))]);

%visualize joint distribution
if i==1
    ylabel('Analytical','FontWeight','bold');
end
if i==3
    yyaxis right
    set(gca,'ytick',[])
    ylabel('mRNA');
    set(gca,'ytick',[],'ycolor','k');
end
xlabel('pre-mRNA');
end

set(figure(1),'Position',[254 561 824 319]);
set(figure(2),'Position',[254 561 824 319]);
set(figure(3),'Position',[402 316 654 606]);
return

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