summarize selected probes

May 13, 2020

0.0.1 Summarize selected probes

- Get the probes selected in Dmel and add properties from current version of the probe designer
- Also output a fasta file to use for blasting against the Dmel transcriptome

```
[1]: #Imports
     import sys
     import pandas as pd
     import os
     import primer3
     from Bio.SeqUtils import MeltingTemp as mt
     import primer3
     from Bio.Seq import Seq
     sys.path.append('../scripts/')
     from plot_helpers import *
     sys.path.append(os.path.join(probe_designer_dir, 'scripts'))
     import screen_kmers
     import choose_probes
     %matplotlib inline
     %load_ext autoreload
     %autoreload 2
```

```
[2]: outdir = '../figures/F1/'
os.makedirs(outdir, exist_ok = True)
```

```
[3]: #Get the probe sequences, and add to the df

qpcr_dir = os.path.join(results_dir, 'qPCR_data')

probe_seqs = os.path.join(qpcr_dir, 'probe_seqs.csv')

seq_df = pd.read_csv(probe_seqs, index_col = 'probe_name')

#Now get the properties for the probes selected for Drosophila

#chosen probes, these are the ones that were included in the Ribo-Pop mix for_

→sequencing

chosen_probes_18S = [12, 18, 21, 24, 28]

chosen_probes_28S = [36, 37, 38, 39, 40, 41, 42, 43, 44, 45]

chosen_18S = seq_df.loc[seq_df['probe_num'].isin(chosen_probes_18S),_□

→['sequence']].copy()
```

```
chosen_28S = seq_df.loc[seq_df['probe_num'].isin(chosen_probes_28S),__
     → ['sequence']].copy()
    chosen 18S['target name'] = '18S'
    chosen 28S['target name'] = '28S'
    chosen_df = pd.concat([chosen_18S, chosen_28S])
    chosen df.reset index(drop = True, inplace = True)
    chosen_df['probe_num'] = chosen_df.index + 1
    chosen_df['length'] = chosen_df['sequence'].apply(lambda x: len(x))
    chosen_df['unique_id'] = chosen_df.apply(lambda x: '%s_%s' % (x['target_name'],_
     \rightarrowx['probe_num']), axis = 1)
    chosen_df.set_index('unique id', drop = False, inplace = True)
[4]: | #Get the values calculated from the probe design pipeline and add
    dmel_18S = os.path.join(results_dir, 'probe_design_results/dmel_200504/
     →probe_design/18S/potential_probes_filt.csv')
    dmel 28S = os.path.join(results dir, 'probe design results/dmel 200504/

→probe_design/28S/potential_probes_filt.csv')
    dmel 18S df = pd.read csv(dmel 18S)
    dmel_28S_df = pd.read_csv(dmel_28S)
    allfilt_df = pd.concat([dmel_18S_df, dmel_28S_df])
    cols2write = ['Tm','sequence', 'target_start', 'target_end','passed_excluded',_
     → 'hairpin_dG', 'homodimer_dG', 'passed_structure',
                 'GC content', 'A content', 'C content', 'GC content rule',
     →'A_composition_rule', 'C_composition_rule',
                 '4xA_stack_rule', '4xC_stack_rule', 'earlyCs_rule', 'any5_rule',
     col_order = ['probe_num', 'sequence', 'target_name', 'target_start', __
     'C_content', 'rolling_Tm_quantile_co', 'hairpin_dG', 'homodimer_dG', |
     'A_composition_rule', 'C_composition_rule', '4xA_stack_rule', '4xC_stack_rule', _
     annotated_df = pd.merge(chosen_df, allfilt_df[cols2write], left_on = __
     annotated df.set index('unique id', inplace = True)
    annotated_df[['dimer_dG', 'dimer_partner']] = choose_probes.
     →calc_dimer(annotated_df)
    annotated_df.reset_index(inplace = True)
```

annotated_df[col_order].round(2).to_csv(os.path.join(outdir,_

→'Dmel_selected_properties.csv'), index = False)

```
[5]: #write fasta file of the probes -- note that we expect matches to align to the

inegative strand

with open(os.path.join(outdir, 'Dmel_probes.fa'), 'w') as g:

for i in annotated_df.itertuples():

g.write('>%s\n%s\n' % (i.unique_id, i.sequence))
```