# **Open Access** Computational methods to identify novel methyltransferases Tanya C Petrossian and Steven G Clarke

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#### Background

1.2% of the yeast genes are estimated to encode enzymes that catalyze the transfer of a methyl group from S-adenosylmethionine (AdoMet) to protein, nucleic acid, lipid, and small molecule substrates [1]. These enzymes function in biosynthesis, regulating metabolic pathways, and controlling gene expression, including writing the histone code. BLAST and MEME/MAST analysis using the amino acid sequence of motifs have previously generated a list of putative Class I methyltransferases [2]. Recently we have used a combination of a new search algorithm and structural information to refine this analysis [3]. This study utilizes these updated methods of identifying motifs and scanning the proteome to predict new members of the different families of methyltransferases in different organisms. These new members may function in novel pathways or new modes of regulation.

#### Materials and methods

Advanced hidden Markov models (HMM) profiles, predicted secondary structures, and solved crystal structures are used to identify the AdoMet-binding motifs of the different families of methyltransferases [1,3]. To generate a list of putative methyltransferases, we used both our newly developed program "Multiple Motif Scanning" [3,4] and HHpred [5]. Sequence similarity networks are then used to predict the probable substrates for the putative methyltransferases [3]. Additionally, several of the candidate methyltransferases were incubated with radioactive Ado-Met to reveal binding by detection of the radioactive protein-ligand via SDS-PAGE separation [1].

### Conclusion

The putative list of methyltransferases for S. cerevisiae among four of the methyltransferases families are italicized (see Table 1). Known methyltransferases are shown for only the SET and SPOUT families. Several putative methyltransferases are found to bind AdoMet through UV-crosslinking experiments (designated \* in Table 1). This approach validated previously suggested putative enzymes and additionally identified several new candidates [3]. Extending this analysis to the human proteome surprisingly reveals little expansion of family members (Figure 1). Our goal is to enhance the functional identification of novel methyltransferases by providing lists of the best candidates for biochemical analyses.

Table I: Proteins classified into four families of methyltransferases

Seven-Beta Strand (Class I) (Not shown here are 33 known species)		SET	SPOUT	N6-Adenosine
YBR141C	YLR137W	Set l	Trm10	Ime4
YBR225W	YMR209C*	Set2	Mrml	Kar4
YBR261C*	YMR228W	Set3	Trm3	YGR001C
YBR271W	YNL022C	Set4	Emgl	
YDR316W	YNL024C	Set5	YGR283C	
YHR209W*	YNL092W	Set6	YMR310C	
YIL064W	YOR239W	Rkml	YOR021C	
YILI I OW		Rkm2		
YJR I 29C*		Rkm3		
YKL155C*		Rkm4		
YKL162C		Ctm I		
YLR063W				YHL039W

	Yeast	Human
Total	~6600 protein- coding genes	~23000 protein- coding genes
Precorrin-like	2 proteins	1 protein
Membrane	3 proteins	3 proteins
Homocysteine	2 proteins	3 proteins
N6-Adenosine	3 proteins	5 proteins
MetH Activation	0 proteins	1 protein
Radical SAM	0 proteins	4 proteins
SET	12 proteins	~56 proteins

#### Figure I

Comparison of the number of known and putative yeast and human methyltransferases in several families.

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#### References

- Petrossian TC and Clarke SG: Bioinformatic identification of Т novel methyltransferases. Epigenomics 2009 in press. Katz JE, Dlakić M and Clarke S: Automated identification of
- 2. putative methyltransferaess from genomic open reading frames. Mol Cell Proteomics 2003, 2:525–540. Petrossian TC and Clarke SG: Multiple Motif Scanning to
- 3. identify methyltransferases from the yeast proteome. Mol. Cell. Proteomics 2009, 8:1516-1526.
- Multiple Motif Scanning. http://www.chem.ucla.edu/files/Motif-4.
- Setup.Zip. Söding J, Biegert A and Lupas AN: The HHpred interactive server for protein homology detection and structure prediction. Nucleic Acids Res 2005, 33:W244–W248. 5.

