Transgenesis of mammalian PABP reveals mRNA polyadenylation as a general stress response mechanism in bacteria

Highlights

- Trans expression of mammalian PABPN1 stabilizes polyadenylated mRNAs in E. coli.
- PABPN1 expression phenocopies pcnB mutation and regulates plasmid copy number.
- 3' polyadenylation acts as a general stress response mechanism in bacteria.
- This study indicates an evolutionary significance of PABP in mRNA metabolism.

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Transgenesis of mammalian PABP reveals mRNA polyadenylation as a general stress response mechanism in bacteria

Nimmy Francis¹,² and Rakesh S. Laishram¹,³,*

SUMMARY
In eukaryotes, mRNA 3'-polyadenylation triggers poly(A) binding protein (PABP) recruitment and stabilization. In stark contrast, polyadenylation marks mRNAs for degradation in bacteria. To study this difference, we trans-express the mammalian nuclear PABPN1 chromosomally and extra-chromosomally in Escherichia coli. Expression of PABPN1 but not the mutant PABPN1 stabilizes polyadenylated mRNAs and improves their half-lives. In the presence of PABPN1, 3'-exonuclease PNPase is not detected on PA-tailed mRNAs compromising the degradation. We show that PABPN1 trans-expression phenocopies pcnB (that encodes poly(A) polymerase, PAPI) mutation and regulates plasmid copy number. Genome-wide RNA-seq analysis shows a general up-regulation of polyadenylated mRNAs on PABPN1 expression, the largest subset of which are those involved in general stress response. However, major global stress regulators are unaffected on PABPN1 expression. Concomitantly, PABPN1 expression or pcnB mutation imparts cellular tolerance to multiple stresses. This study establishes mRNA 3'-polyadenylation as a general stress response mechanism in E. coli.

INTRODUCTION
Polyadenylation (addition of a poly-adenosine tail, PA-tail) by poly(A) polymerase (PAP) enzymes at the 3'-end is a key-processing event of nascent transcripts that determines fate of a cellular messenger RNA (mRNA) (Bardwell et al., 1990; Dreyfus and Regnier, 2002; Shatkin and Manley, 2000; Zheng and Tian, 2014). In mammals, almost all mRNAs are polyadenylated (~250 adenosines) inside the nucleus, which are subsequently shortened (~70–100 adenosines) after cytoplasmic processing (Eckmann et al., 2011; Jalkanen et al., 2014; Laishram, 2014). PA-tail endows stability to the mRNA and is required for efficient translation (Glaunsinger and Lee, 2010; Laishram, 2014). On the contrary, PA-tail addition marks mRNAs for degradation in prokaryotes (Belasco, 2010; Hajnsdorf et al., 1995; Hajnsdorf and Regnier, 1999; Sarkar, 1997). Also, PA-tails are short (15–40 adenosines in E. coli) and are present on limited mRNAs (Sarkar, 1997). However, recent studies indicate widespread polyadenylation of bacterial transcripts including ribosomal RNAs, transfer RNAs, non-coding RNAs, small RNAs and mRNAs (Li et al., 1998; Maes et al., 2016; Mohanty et al., 2012; Reichenbach et al., 2008; Xu et al., 1993). Polyadenylated transcripts largely have structured 3'-ends, and that addition of a PA-tail provides platform for the 3'-exonuclease polynucleotide phosphorylase (PNPase) to initiate 3'-S' exonucleolytic degradation along with other degradosome complex components [ribonuclease E (RNase E), RNA helicase B (RhlB) and enolase] (Blum et al., 1999; Carposis, 2007). Thus, bacterial polyadenylation primarily regulate turnover and quality control of specific cellular transcripts (Mohanty and Kushner, 2011).

Poly(A) polymerase I (PAPI) encoded by pcnB gene is the primary PAP in E. coli (Liu and Parkinson, 1989; Xu et al., 1993), yet, pcnB gene is dispensable in the cell (Masters et al., 1993). On the other hand, PAPI overexpression exhibits cellular toxicity by rapid polyadenylation of tRNAs and inhibiting protein synthesis (Mohanty and Kushner, 2012). Therefore, PAPI level is maintained low in vivo via a non-canonical initiation codon and a poor ribosome binding site despite transcribing from multiple promoters (three e70-and two sS-dependent promoters) (Bims and Masters, 2002; Jasiecki and Wegrzyn, 2006; Nadratowska-Wesołowska et al., 2010). PAPI is primarily known for the regulation of colE1-based plasmid replication by controlling RNAI transcript stability (He et al., 1993; Xu et al., 1993). Recent studies have now established role of PAPI-mediated polyadenylation in functional gene expression and have linked with stationary phase...
growth, chemotaxis and motility, nutrient starvation, and envelop stress (Aiso et al., 2005; Carabetta et al., 2010; Joanny et al., 2007; Maes et al., 2013, 2016; Mohanty et al., 2012, 2020; Reichenbach et al., 2008; Sinha et al., 2018). However, the complete pool of polyadenylated mRNAs and a wide range of physiological processes regulated by general polyadenylation are yet to be determined.

Poly(A) binding proteins (PABPs) bind PA-tail and protects PA-tailed mRNAs from nuclease degradation in eukaryotes (Bernstein and Ross, 1989; Kuhn and Wahle, 2004). In mammals, there are three cytoplasmic PABPs (PABPC1, PABPC2, and PABPC3) and one nuclear PABP (PABPN1) (Kuhn and Wahle, 2004). While cytoplasmic PABP is critical for the stability and translation of the mRNA, nuclear PABPN1 primarily promotes stability to the PA-tailed mRNAs and regulates PA-tail length control and mRNA nuclear export (Lemay et al., 2010). Two point mutations in the RNA binding domain of PABPN1 (tyrosine 175 to alanine, Y175A and phenylalanine 215 to alanine, F215A) are shown to compromise PA-tail mRNA binding and PABPN1 function (Kerwitz et al., 2003; Kuhn et al., 2003). No counterparts of eukaryotic PABPN1 that protects and stabilizes cellular PA-tailed mRNAs are known in bacteria (Dreyfus and Regnier, 2002). Two RNA binding proteins Hfq and cold shock-like protein E (CspE) in E. coli are reported to bind PA-containing RNAs in vitro (Feng et al., 2001; Folichon et al., 2003). Hfq can also promote PA-tail elongation by PAPI (Hajnsdorf and Regnier, 2000; Mohanty et al., 2004). Nonetheless, both CspE and Hfq interact with multiple RNA sequences and their effect on the PA-tail stabilization in vivo is unclear (Hajnsdorf and Boni, 2012; Hajnsdorf and Regnier, 2000; Mohanty et al., 2004; Phadtare and Inouye, 1999). Therefore, cellular functions of CspE and Hfq on mRNA metabolism are different from that of eukaryotic PABP mediated mRNA stabilization.

To understand the difference in the cellular roles of polyadenylation on mRNA metabolism in bacteria and eukaryotes, we trans-express mammalian PABPN1 in E. coli. Expression of PABPN1 but not the mutant PABPN1 defective for PA-tail binding stabilizes earlier reported polyadenylated mRNAs in E. coli. Consistent mRNA stabilization was observed in pcnB mutation with no additional effect of PABPN1 expression suggesting that PABPN1 affects polyadenylated transcripts. PABPN1 trans-expression also phenocopies pcnB mutation and stabilizes RNA transcript thereby reducing copy number of colE1-based plasmids. We demonstrate that PABPN1 trans-expression compromises PNPase association and degradation of polyadenylated mRNAs. Genome-wide RNA-Seq analysis shows up regulation of a large number of transcripts on PABPN1 expression in E. coli. Functional analysis of the up regulated mRNAs reveals involvement of polyadenylation in new cellular functions that include stress response, membrane transport, cellular metabolism, xenobiotic degradation, and pathogenesis. Among the upregulated mRNAs, the largest fraction (~33%) is involved in response to multiple stresses such as osmotic, oxidative, acid, DNA alkylation, heat and cold stresses but without affecting the expression of known stress response regulators. We demonstrate that pcnB mutation and introduction of PABPN1 but not the mutant PABPN1 (compromised for PA-tail binding) stimulates expression of stress response genes and imparts cellular tolerance to multiple stresses. Our study establishes 3’-end mRNA processing as a general stress response mechanism in bacteria.

RESULTS

Trans-expression of mammalian nuclear poly(A) binding protein, PABPN1 in E. coli

In eukaryotes, PAB binding primarily protects PA-tails at the mRNA 3’-end and stabilizes polyadenylated mRNAs. Therefore, to assess the effect of PABP on mRNA turnover in bacteria, we trans-expressed mammalian nuclear PABPN1 in E. coli K-12 strain MG1655 and its isogenic pcnB mutant strain (Figure 1A, Table 1). PABPN1 was expressed first through pBAD18 plasmid and PABPN1 expression was tested by Western blot analysis using PABPN1-specific antibody. We observed PABPN1 expression in the pBAD18-PABP transformed cells but not in the plasmid control pBAD18 transformed cells after arabinose induction (Figure 1B). Near similar PABPN1 expression was observed in all media after induction with 0.5% arabinose (LB, Minimal-Glucose, or Minimal-Glycerol) in strains transformed with pBAD18-PABP but not in pBAD18 plasmid (Figure S1A). In a second approach, we expressed PABPN1 stably through chromosomal insertion using λ-Inch technique (Boyd et al., 2000) to obtain a stable single copy expression (Figure 1A). We confirmed PABPN1 integration into the genomic DNA by PCR analysis and expression by Western blot analysis in both wild type and pcnB mutant cells (Figures 1C and S1B). Hereafter, stable PABPN1 expressing cells are referred to as MG-PABP or pcnB-PABP respectively in the text. We observed similar results from both plasmid-based PABPN1 expression and chromosomal PABPN1 expression.
Transgenesis of mammalian PABPN1 stabilizes polyadenylated mRNAs in E. coli

qRT-PCR analysis demonstrated that expression of PABPN1 increased levels of earlier known polyadenylated transcripts (lpp, secG, rpsO, cspE, trpA, osmY, and RNAI) (Mohanty and Kushner, 2011) but not control non-polyadenylated transcript dxs (Figure 1D). Strikingly, there was >3- to 8-fold increase in the expression levels of transcripts tested on PABPN1 expression. Introduction of a PA-tail binding mutant (PABP-Y175A) compromised the stabilization of mRNAs studied (Figure 1D). Consistently, expression levels of the same transcripts were also enhanced on pcnB mutation (Figure 1D). We also confirmed polyadenylation of one of the transcript (osmY) by 30-RACE assay followed by 30-end PA-tail sequencing (Figures 1E and1 F). 30-RACE assay also showed an increased level of polyadenylated osmY transcript and a concomitantly higher total osmY mRNA level on PABPN1 expression (Figures 1D, F, S1C). While pcnB mutation resulted in the loss of 30-RACE product, it still augmented the expression level of osmY mRNA similar to that of PABPN1 expression (Figures 1D and 1F). Further, measurement of mRNA stability after inhibition of transcription with rifampicin treatment showed around 2- to 4-fold increase in the half-life (T 1/2) of known polyadenylated mRNAs (osmY, RNAI) on PABPN1 (but not PABPN1-Y175A mutant) expression (Figure 2A).
Similar enhancement in the half-life was also visible in the case of pcnB mutation. However, there was no additional induction of half-life or mRNA levels on PABPN1 expression in the pcnB mutant background (Figure 2A). Together, these results indicate that PABPN1 stabilizes polyadenylated transcripts in E. coli.

We then tested PABPN1 binding to one of the bacterial polyadenylated transcripts in vitro. We synthesized a radiolabeled RNAI with PA-tail (16 adenosines, RNAI-A16) and without PA-tail (non PA-tail, RNAI-A0). Subsequently, in EMSA experiments, we observed purified recombinant PABPN1 binding to the RNAI-A16 but not to the non PA-tail RNAI-A0 (Figures 2B and S1D). The binding was abolished by introduction of a PA-tail binding mutation on PABPN1 (Y175A) (Figure 2B). RNA immunoprecipitation (RIP) experiment also indicated PABPN1 association on the known polyadenylated transcripts osmY, secG and rpsO mRNAs in the cell (Figure 3I). On the other hand, bacterial adenosine binding protein CspE exhibited little affinity toward RNAI-A16 or RNAI-A0, whereas Hfq was bound to both the transcripts (Figure S1 E–S1G). Moreover, ectopic expression of CspE or Hfq had no effect on the expression levels of known PA-tailed mRNAs (lpp, secG and RNAI) and the plasmid copy number unlike PABPN1 expression (Figure S1H and S2A) (PABP expression and plasmid copy number control is detailed in the following section). Consequently, pcnB mutation but not cspE mutation stabilized PA-tailed mRNAs (rpsO, lpp, secG, RNAI, trpA) (Figure S2B). However, hfq mutation showed a moderate enhancement in the mRNA levels (Figure S2B) likely through its effect on PAPI activity as described earlier (Hajnsdorf and Regnier, 2000). These results indicate that known E. coli adenosine-binding proteins are functionally distinct from PABPN1 in terms of PA-tailed mRNA metabolism and that PABPN1 binding specifically stabilizes polyadenylated transcripts in E. coli.

**PABP expression phenocopies pcnB mutation, alters plasmid copy number control, and compromises PNPassociation**

Since polyadenylation is known to regulate colE1-based plasmid replication through RNAI transcript stability (Xu et al., 1993), we analyzed plasmid copy number control of colE1-based plasmids after PABPN1
expression. While MG1655 cells grew till 2000-μg/mL ampicillin, PABPN1 expressed cells but not mutant PABPN1 (Y175A) expressed cells were sensitive at 500-μg/mL ampicillin when transformed with pBAD18 plasmid (Figure 2C). Control pcnB mutant cells also showed sensitivity at 500-μg/mL ampicillin indicating a decrease in the plasmid copy number on PABPN1 expression or pcnB mutation. Similar sensitivity was also observed in dilution plating of the same strains at 500 and 1000-μg/mL ampicillin (Figures 2D and S2C). Subsequently, pBAD18 plasmid content was reduced in control pcnB mutant and PABPN1 expressed

Figure 2. Trans-PABPN1 expression phenocopies pcnB mutation and alters plasmid copy number control

(A) Half-life (T½) measurement of cellular polyadenylated transcripts, RNAI and osmY, and a non-polyadenylated control transcript, dks after inhibition of transcription with rifampcin from various strains as indicated. T½ is expressed in minutes (min). Data are mean ± SEM of n = 3 independent experiments.

(B) RNA EMSA experiments carried out with in vitro transcribed radiolabeled RNAI transcript with a PA-tail (RNAI-Au) and no PA-tail (RNAI-A0) in the presence of increasing recombinant His-PABPN1 or PA-tail binding mutant His-PABPN1 Y175A (PABPM) (0–30 nM). RNAI-PABPN1 binary complex, B and unbound RNAI, F and free RNAI probe, FP, are indicated.

(C) Antibiotic sensitivity assay of wild-type MG1655, PABPN1 expressed in MG1655 (MG-PABP) or pcnB mutant (MG- pcnB) and PABPN1 expressed in pcnB mutant (pcnB-PABP) cells transformed with pBAD18 plasmid and streaked on plates supplemented with increasing ampicillin concentration from 100 to 2000 μg/ml as indicated.

(D) Dilution spotting of various strains as in “C” from 10⁻¹ to 10⁻⁶ dilutions on LB plate supplemented with 100 and 1,000 μg/mL ampicillin as indicated.

(E) Analysis of plasmid content of pBAD18 and control pCL1920 plasmids isolated from various strains as in “C” after digestion with NdeI to linearize both plasmids.

(F) Copy numbers of pBAD18 and pCL1920 plasmid DNA per 1 OD₆₀₀/ml (x10⁹) from various strains as in “C” as indicated. See also Figures S1 and S2.
both chromosomally expressed and plasmid expressed) cells with no effect on control RNAi-independent plasmid pCL1920 (Figures 2E and S2D). There was ~4-fold reduction in the copy number per OD_{600} per ml of pBAD18 plasmid on pcnB mutation and PABPN1 expression (Figure 2F). Consistently, there was increased stability of RNAi transcript in PABPN1-expressed cells and pcnB mutant cells with no effect on control bla expression from pBAD18 or aad1 mRNA expression from pCL1920 plasmid (Figure 3A). These
results indicate that PABPN1 expression phenocopies *pcnB* mutation and that PABPN1 expression alters plasmid copy number.

To further confirm the stabilization of polyadenylated RNAI, we employed in vitro transcribed radiolabeled RNAI-A<sub>16</sub> and RNAI-A<sub>0</sub> transcripts and tested in active MG1655 cell lysate in the presence and absence of PABPN1 expression. As expected, RNAI-A<sub>16</sub> was degraded more rapidly than RNAI-A<sub>0</sub> in active cell lysate (Figure 3B). We also observed stabilization of RNAI-A<sub>16</sub> but not RNAI-A<sub>0</sub> in the presence of PABPN1 expression (Figures 3C and 3D). Strikingly, RNAI-A<sub>16</sub> that was degraded after ~2 min in active cell lysate was stabilized by PABPN1 but not mutant PABPN1-Y175A expression (Figure 3C). However, RNAI-A<sub>0</sub> decay was not affected by PABPN1 expression (Figure 3D). A control mammalian heme oxygenase 1 (HMOX1) UTR RNA was degraded alike in both MG1655 and PABPN1 expressed cell lysates (Figure 3E) indicating similar ribonuclease activities of the lysates and the degradation of mRNA. We also observed increased stabilization of RNAI-A<sub>16</sub> in the presence of increasing amounts of recombinant PABPN1 (Figure S2E). To further understand how PABPN1 binding of the PA-tail stabilizes polyadenylated mRNAs in *E. coli*, we pulled down proteins bound on an in vitro transcribed biotinylated RNAI-A<sub>16</sub> from cell-extracts of wild type MG1655 and PABPN1 expressed cells and analyzed for the 3′-exonuclease PNPase association (Figures 3F and 3G). We detected PNPase on RNAI-A<sub>16</sub> transcript from wild type cell lysate (Figure 3G). However, PNPase was not detected on RNAI-A<sub>16</sub> from the PABPN1 expressed cell lysate (Figure 3G). Consistently, RIP experiments showed PNPase association with polyadenylated transcripts (secG, osmY, and rpsO) in MG1655 but not in PABPN1 expressed cells (Figures 3H and 3I). Control RNA polymerase (RNP) was detected on all the transcripts in both cell types. However, PABPN1 was specifically detected on transcripts from the PABPN1 expressed cells and not from the wild type cell lysate (Figure 3I). Together these results indicate that PABPN1 expression compromises PNPase-mediated 3′-exonucleolytic degradation of polyadenylated mRNAs.

**PABPN1 expression alters global landscape of mRNA polyadenylation in *E. coli***

Since PABPN1 expression stabilizes known polyadenylated mRNAs in *E. coli*, we carried out genome-wide RNA-Seq analysis of wild-type and PABPN1 expressed (both plasmid-based and chromosomally expressed) MG1655 cells to assess the extent of mRNA polyadenylation in the cell. Of approximately 4,200 protein coding mRNAs in *E. coli* (Serres et al., 2001), we detected more than 3,800 mRNAs (~90% of the total genes) in our RNA-Seq analysis (Figures 4A and 4B). Of these transcripts, we observed approximately 700 mRNAs upregulated on both chromosomal-based PABPN1 expression and plasmid-based PABPN1 expression (Figures 4A and 4B). A list of mRNAs with altered expression from both plasmid and chromosomal PABPN1 expression is shown in Table S1. More than 75% of the upregulated genes were common between the plasmid-based and chromosomal PABPN1 expressed cells (Figure 4C). A list of commonly upregulated mRNAs is shown in Table S2. We then validated 10 genes (*aidB, aldB, bsmA, clpA, dsdA, osmY, otsA, uspA, uspC, uspE, and wrbA*) from our RNA-Seq by quantitative real-time PCR (qRT-PCR) analysis (Figure 4D) and 2 mRNAs (osmY and clpA) by 3′-RACE assay (Figure S3A). We observed up regulation of these mRNAs on PABPN1 expression or *pcnB* mutation in qRT-PCR analysis (Figure 4D). There was similar induction in the 3′-RACE product on PABPN1 expression that was diminished on *pcnB* mutation (Figure S3A).

Further, to validate the induced polyadenylated mRNA level on PABPN1 expression in the cell observed in 3′-RACE, we isolated total cellular PA-plus RNA and analyzed polyadenylated mRNAs by a 3′-end G-I tailing coupled PA-tail assay (Figure S2F) (Kusov et al., 2001; Patil et al., 2014). Here, a short stretch of guanosine and inosine (G-I) nucleotides were added on the *in vivo* isolated PA-plus RNA using yeast PAP (Kusov et al., 2001). Specific mRNAs were further amplified using a reverse primer specific to the added G-I tail and a gene-specific forward primer as in the case of 3′-RACE assay, or by qRT-PCR analysis using the similar primers for quantification (Figure S2F). We consistently observed increased polyadenylated mRNA level (*lpp, secG, rpoS, cspE, uspA, uspE, wrbA, rmf, cspD, and osmY*) in the presence of PABPN1 expression in the cell while non-polyadenylated control dks was negligibly detected in both MG1655 and MG-PABP cells (Figure S3B). Similar inductions in the expression levels of these mRNAs (*lpp, secG, rpoS, cspE, trpA, uspA, uspC, uspE, cspA, wrbA, rmf, cspD, otsA, and osmY*) were also seen in the qRT-PCR analysis of PA-plus mRNAs (Figure S3C). In line with this, we also observed increased levels of known polyadenylated non-coding RNAs including sRNAs (GlmY, SroH, GlmY) and tRNAs (valV, pheU, hisR, cysT) (Mohanty et al., 2020) on PABPN1 expression from the PA-plus RNA analysis (Figure S2D and S3E). However, while majority (~90%) of mRNAs were detected in our RNA-Seq, small RNAs (tRNA or sRNA) were minimally...
Figure 4. Genome-wide RNA-Seq reveals a new polyadenylation-mediated stress response pathway in *E. coli*

(A) Heatmap showing altered gene expression profile from plasmid-based PABPN1 expressed (plasmid-PABP) and chromosomally expressed PABPN1 (chromosomal PABP) in MG1655 cells compared with the wild-type MG1655 cells.

(B) Table showing number of transcripts altered both upregulated and downregulated on both plasmid PABPN1 expression and chromosomal PABPN1 expression. Complete list of transcripts altered on PABPN1 expression is shown in Table S1.

(C and D) Venn diagram showing upregulated genes overlapped between PABPN1 expressed chromosomally and PABPN1 expressed through plasmid (D) qRT-PCR analysis of 10 select polyadenylated mRNAs from the RNA-Seq data using total RNA isolated from MG1655, *pcnB* mutant (MG-*pcnB*), PABPN1 (MG-PABP) expressed cells as indicated (p values for all the genes were <0.001).

(E) Functional pathway analysis of upregulated genes common between PABPN1 expressed chromosomally and PABPN1 expressed through plasmid showing polyadenylated mRNAs involved in multiple cellular functions. Complete list of commonly upregulated mRNAs is shown in Table S2.

(F) List of select genes involved in various stress response pathways from the commonly upregulated genes between PABPN1 expressed chromosomally and PABPN1 expressed through plasmid. Some of the global regulators of stress response that are not affected by PABPN1 expression are shown in the right.

(G) qRT-PCR analysis of *rpoS*, *oxyR* and target gene *bsmA* using total RNA isolated from MG1655, *pcnB* mutant (MG-*pcnB*), PABPN1 (MG-PABP) expressed cells as indicated (p values for all the genes were <0.001).

See also Figures S3 and S4.
detected in our RNA-Seq (18 of ~86 total tRNA genes and >85 total sRNAs in *E. coli*). Therefore, we screened total cellular sRNAs and tRNAs using the G-I tailing coupled PA-tail assay described above to assess the effect of PABPN1 trans-expression on these RNAs (Figure S2F). Our screening identified 74 tRNAs and 72 sRNAs, of which RNA levels of only ~19 sRNAs and a majority of ~40 tRNAs were induced on PABPN1 expression. The total list of sRNAs and tRNAs that were stimulated or unchanged on PABPN1 expression from the screening are shown in Table S3. PA-tail assay coupled qRT-PCR screening of 42 select tRNAs and 42 sRNA, along with gel analysis of ~22 select sRNAs and tRNAs from the PA-tail assay is shown in Figures S4A–S4D. Concomitantly, we also observed PABPN1 association with the sRNAs and tRNAs that were induced on PABPN1 expression but not with the unaffected RNAs in the cell (Figures S2F–S2H).

**mRNA polyadenylation as a novel stress response mechanism in bacteria**

To understand the cellular functions affected by PABPN1 expression, we carried out functional pathway analysis of the upregulated genes on PABPN1 expression (Figure 4E). For our analysis, we considered only those mRNAs that were upregulated in both plasmid-based and chromosomal expression of PABPN1. Our functional analysis revealed involvement of polyadenylation in new cellular functions including membrane transport, different metabolic pathways, pathogenesis, xenobiotic degradation, maintenance of cell wall and membrane integrity, antimicrobial resistance, and aerobic respiration (Figure 4E). Among the PA-tailed mRNAs, the largest fraction (>33%) were those involved in a wide range of stress response pathways including DNA damage, oxidative stress, osmotic stress, heat or cold, nutrient starvation, or biofilm formation (Figures 4E and 4F; Table S4). However, expression of many of the global regulators of stress response including the master regulator gene of stress response *rpoS* (Battesti et al., 2011; Gottesman, 2019; Starosta et al., 2014) or stress regulator *oxyR* was unaffected (Figure 4G). Whereas, known targets of *rpoS* or *oxyR* were up regulated on PABPN1 trans-expression (Figure 4G). Moreover, tRNA polyadenylation while affects functional tRNA levels, it primarily regulates cellular translation (Mohanty and Kushner, 2012; Mohanty et al., 2012). Furthermore, majority of the stress responsive sRNAs (e.g. OxyS, OmrA, MicC, MicF, GcvB, MgrR, OmrB, RybB, FnrS, ArcZ, DsrA, RprA, and DicF) were unaffected by PABPN1 expression (Table S3) (Gottesman and Storz, 2011; Hobbs et al., 2010; Holmquist and Wagner, 2017). Together, these results indicate a direct PA-tail stabilization of stress response mRNAs by PABPN1 trans-expression and reveal a new role of global mRNA polyadenylation in general stress response pathway.

**PABPN1 expression or *pcnB* mutation alters sensitivity of *E. coli* cells to multiple stresses**

To investigate the role of polyadenylation-mediated mRNA turnover in bacterial stress response, we tested stress sensitivity of wild type, *pcnB* mutant, and PABPN1 or PABPN1 (Y175A) PA-tail binding mutant expressed MG1655 cells in six different stress conditions (osmotic stress, oxidative stress, DNA damage/alkylation, acid shock, heat shock and cold shock). While MG1655 cells were viable till $10^{-6}$ dilutions in control condition, their growth was compromised in all stresses, which was substantially ameliorated by *pcnB* mutation or PABPN1 expression but not by mutant PABP-Y175A expression (Figure 5A). Viable colonies (cfu/mL) were also reduced (~5-10-fold) in wild-type and mutant PABP-Y175A expressed cells with no significant effect in *pcnB* mutant or PABPN1 expressed cells in different stress conditions (Figure 5B). Likewise, in growth curve analysis, all cells reached OD$_{600}$ of ~3.0 under control condition in 12 h, wild-type cells reached OD$_{600}$ only ~1.0 on stress treatment (NaCl, MNNG and acid) (Figure 5C, 5D, and 5E). *pcnB* mutation or PABP expression resulted in a considerable recovery in the growth to ~2.1 OD$_{600}$ under the same stresses (Figures 5C and 5D). Moreover, wild type cells exhibited weak biofilm-forming ability compared to *pcnB* mutant and PABPN1 expressed cells (Figure 5F). Further, we also observed a decrease (~20~40%) of wild type cell size under high osmolarity as reported earlier (Dai and Zhu, 2018) but no marked changes on the sizes of *pcnB* mutant and PABPN1 expressed cells (Figures 5E and 5F). However, treatment with DNA alkylating agent MNNG resulted in an elongation (~5-fold) of the wild type cells (El-Hajj and Newman, 2015; Uphoff et al., 2016) but did not markedly affect *pcnB* mutant or PABPN1 expressed cells (Figures 5C and 5D). Consistently, qRT-PCR analysis revealed stimulated expression levels of stress response genes (ranging from >10-fold for *bsmA* or *aiba* to 3- to 5-fold for *clpA*, *cspE*, *otsA*, *osmY*, *aibD*, *deoA*) under different stresses (oxidative stress, osmotic stress and alkylation stress) and on PABPN1 expression (Figure S5E). Western blot analysis also showed induced protein levels of stress response proteins (UspF and DsdA) on stress treatment in the presence and absence of *pcnB* mutation or PABPN1 expression (Figure 5G). We also show that PAPI activity is unaffected by PABPN1 expression and PABP and PAPI are functionally distinct (Figures S5A and S6B). Together, our results from PABPN1 transgenesis reveal that 3'end polyadenylation-mediated mRNA turnover regulates general stress response pathway in *E. coli*. 
DISCUSSION

Polyadenylation at the structured 3′-end is a key post-transcriptional modification of mRNAs in bacteria that initiates 3′–5′ exonucleolytic degradation, a process critical for mRNA turnover and quality control (Blum et al., 1999; O’Hara et al., 1995). 3′-polyadenylation is primarily carried out by PAPI enzyme in E. coli (Cao and Sarkar, 1992). Another protein, the 3′-exonuclease PNPase is also reported to act as a PAP in E. coli (Kalapos et al., 1994; Mohanty and Kushner, 2000), yet the exact mechanism how PNPase functions with PAPI for polyadenylation of different cellular transcripts is unclear and is likely dependent on the presence of inorganic phosphates in the cell. However, in mammals, different PAPs (canonical PAPα, PAPγ, and
Star-PAP) exhibit target specificity with distinct niche of mRNA targets for polyadenylation (Li et al., 2017). No such reports for specific target niche selection are available for PAPI or PNPase in E. coli. Nevertheless, our genome-wide RNA-Seq analysis after PABPN1 trans-expression shows widespread mRNA polyadenylation in E. coli. Moreover, PABPN1 binding alters ramifications of 3′-end polyadenylation on mRNA metabolism in E. coli that now mimics that of eukaryotes in terms of stabilizing the polyadenylated mRNA. Interestingly, polyadenylation can also mediate RNA decay in eukaryotic cellular organelles such as chloroplast and mitochondria (Schuster and Stern, 2009). While the role of polyadenylation in plant cell chloroplast is similar to that in bacteria, the polyadenylated RNAs in mitochondria are diverse (Levy and Schuster, 2016). Mammalian cell mitochondria harbor both stable polyadenylated mRNAs, and truncated transcripts that are targeted for degradation by polyadenylation (Slomovic et al., 2005). This indicates an intricate evolutionary relationship between polyadenylation and RNA metabolism.

However, unlike in mammals, PABPN1 does not affect PAPI activity or PAPI function (Figure S6A) (Kerwitz et al., 2003). PABPN1 does not exist naturally in E. coli and there is also no bacterial counterpart of PABPN1 that specifically protects cellular mRNA PA-tails (Dreyfus and Regnier, 2002). There are two RNA binding proteins, Hfq and CspE that are shown to bind RNAs containing poly-adenosine track in E. coli. However, they are functionally different from PABPN1 in PA-tailed RNA metabolism, and their stabilizing effect on PA-tailed mRNA in vivo is unclear (Hajnsdorf and Boni, 2012; Hajnsdorf and Regnier, 2000; Mohanty et al., 2004; Phadtare and Inouye, 1999). Thus, this study indicates PABP as a protein at an evolutionary crossroad, emergence of which marks an overturn in the cellular function of polyadenylation on mRNA metabolism (Dreyfus and Regnier, 2002; Hernandez et al., 2010). Our phylogenetic analysis also revealed absence of PABP but presence of PA-tail in archae- and eu-bacteria, and that the PA-tail-mediated mRNA stabilization appeared only in eukaryotes in parallel with the emergence of PABP (Figure S6B).

We have successfully expressed a functional mammalian PABPN1 in E. coli, and that PABPN1 binding stabilizes polyadenylated transcripts such as osmY. While eukaryotic transgene expression in bacteria is a widely employed approach, trans-expression of eukaryotic proteins functional in bacteria that alters bacterial physiology is still limited (Ostermeier et al., 1996). Our study shows a transgenesis that modifies bacterial physiology and gene expression program including that of stress response, membrane transport, metabolite degradation, and biosynthesis pathways. In addition to the laboratory E. coli K-12 strain, the role of polyadenylation in mediating stress response will have ramifications to pathogenic E. coli strains and other gram negative pathogens that have PAPI in the cell (Adilakshmi et al., 2000; Hayashi et al., 2001; Saravanamuthu et al., 2004; Welch et al., 2002). Such bacterial pathogens use different mechanisms to adapt and tolerate severe intracellular stress to survive in a host (Fang et al., 2016). Polyadenylation can be one such mechanism that is used by bacterial pathogens. (Our preliminary work also suggest role of polyadenylation in infection and intracellular survival). Similar transgenesis approach can also be used for genetic and metabolic engineering in E. coli to modulate biosynthetic pathways to facilitate industrial production of secondary metabolites (Pontrelli et al., 2018). This study unveils a new approach employing functional alien trans-gene expression in bacteria and altering the physiology for industrial and medical applications – a progressive step in applied bacteriology.

Our study shows that polyadenylation-mediated mRNA turnover acts as a general stress response mechanism in E. coli and that PABPN1 expression stabilizes stress response mRNAs. Nevertheless, majority of the stress-related regulatory RNAs such as sRNAs are unaffected by PABPN1 expression indicating that PABPN1 primarily stabilizes stress response mRNAs directly through PA-tail binding. We have also confirmed this direct effect of PABPN1 on target PA-tail mRNAs using PABPN1 PA-tail binding mutants. Interestingly, 3′-polyadenylation does not change expression of many of the global regulators of stress response including rpoS gene expression. This indicates that the polyadenylation-mediated pathway is a new and an additional mechanism of stress response. However, it is likely to act cooperatively with the transcriptional- and/or small molecule ppGpp-mediated pathways to generate rapid production of effector stress response proteins (Battesti et al., 2011; Gottesman, 2019). Control of mRNA turnover by polyadenylation will prevent futile cycle of transcriptional stimulation that is countered by degradation pathway during stress. In addition, mRNA turnover rate determines the steady state level of an mRNA in the cell affecting its cellular availability for translation (Nouaille et al., 2017). Our study also reveals that mRNA stabilization during stress induces protein expression of stress response regulators in E. coli.
Limitations of the study

The polyadenylation mediated general stress response pathway identified in the study may be limited to the bacterial system and may not directly apply to stress response in mammalian or human cells. Moreover, the industrial and medical applications of the metabolic engineering deduced from this study require validation with appropriate model organisms. Understanding the role of mRNA turnover through polyadenylation needs further investigation in pathogenic bacterial strains to understand its role in bacterial pathogenesis.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103119.

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AUTHOR CONTRIBUTIONS

N.F. planned and carried out all experiments and R.S.L. analyzed and interpreted the experimental data, as well as conceptualized and wrote the paper.

DECLARATION OF INTERESTS

None declared.
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## STAR METHODS

### KEY RESOURCES TABLE

| Reagent or Resource | Source | Identifier |
|---------------------|--------|------------|
| **Antibodies**       |        |            |
| Mouse monoclonal anti-RNA polymerase β 88B13 antibody | Invitrogen | Cat. No: MA1-25425; RRID: AB_795355 |
| Rabbit monoclonal anti polynucleotide phosphorylase antibody | Bioorbyt | Cat. No: Orb20774 RRID: AB_10936491 |
| Rabbit monoclonal anti-PABPN1 (H-46) antibody | Santacruz | Cat. No: SC-67017 RRID: AB_2156733 |
| Mouse monoclonal β-galactosidase (40-1a) | Santacruz | Cat. No: SC-65670 RRID: AB_831022 |
| Rabbit polyclonal anti-D-serine dehydratase A (DsdA) | Mybiosource | Cat. No: MBS1497575 RRID: NA |
| Rabbit polyclonal anti-universal stress protein F (UspF) | Mybiosource | Cat. No: MBS7000300 RRID: NA |
| **Bacterial and virus strains** | | |
| Please See | Table 1 | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| N-Methyl-N’-Nitro-N-Nitrosoguanidine | Sigma | Cat.No: 129941 |
| Isopropyl Thi-o-β-D-Galactoside | Sigma | Cat.No: 42897 |
| Avidin-Agarose Beads | Thermo Scientific | Cat.No: 20219 |
| biotin16-UTP | Biotium | Cat.No: BU6105H |
| Rifampicin | Himedia | Cat.No: TC-354-5G |
| **Critical commercial assays** | | |
| PA-tail assay kit | Affymetrix | Cat.no: 764551KT |
| **Deposited data** | | |
| All sequencing fastq files generated | This Paper | Accession Number: GSE166974 |
| **Experimental models: Organisms/strains** | | |
| Please See | Table 1 | N/A |
| **Oligonucleotides** | | |
| Please see | Table S5 | N/A |
| **Recombinant DNA** | | |
| Please see | Table1 | N/A |
| **Software and algorithms** | | |
| Image J | NA | https://imagej.nih.gov/ij/download.html |

### RESOURCE AVAILABILITY

#### Lead contact
- Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Rakesh S. Laishram (E mail: laishram@rgcb.res.in).

#### Materials availability
- Plasmids and strains generated in this study are available from the lead contact on request. This study did not generate new unique reagents.
Data and code availability

- The data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO datasets with GEO Series accession number GEO: GSE166974. Accession numbers are listed in the key resources table.

- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

All experiments were carried out in E. coli K-12 strain MG1655 background. Genotypes of strains employed in the study are shown in Table 1. DH5α strain was used for cloning and plasmid amplification while BL21(DE3) was employed for recombinant protein expression. pcnB, hfq, cspE mutant strains were kind gift from Dr. J. Gowrishankar, CDFD, Hyderabad. Mutants from different bacterial backgrounds were transferred to MG1655 by P1 transduction as described previously (Thomason et al., 2007). All bacterial strains were grown at 37°C in nutrient rich, Luria Bertani or Minimal growth media (containing 0.7% potassium phosphate dibasic, 0.2% potassium phosphate monobasic, 0.1% ammonium sulfate, 0.05% sodium citrate, 0.01% 0.02% vitamin B1 and 2 mM magnesium sulfate) supplemented with 0.2% glucose or glycerol respectively (Anupama et al., 2011). Arabinose was supplemented in the medium at 0.5% concentration or unless otherwise indicated in the text. Unless otherwise indicated antibiotics were also supplemented at the concentration of 100 µg/ml for ampicillin, 50 µg/ml for Kanamycin, and 50 µg/ml for spectinomycin as described previously (Anupama et al., 2011).

METHOD DETAILS

Induction of stress response

For stress sensitivity assay, cells were grown to approximately O.D₆₀₀ of 0.5 in LB media. To induce different stress response, bacterial culture was treated with 400 mM NaCl (osmotic shock), 5 mM hydrogen peroxide, H₂O₂ (oxidative stress), 25 µg/ml N-methyl-N’-nitro-N-nitrosoguanidine, MNNG (DNA alkylation stress or DNA damage), low pH at 3.0 adjusted with 100% hydrochloric acid, HCl (acid shock), heat shock at 42°C (heat stress), and cold shock at 4°C (cold stress) for one hour each as described earlier (Rojas et al., 2014). After one-hour, cells were serially diluted from 10⁻² to 10⁻⁸ in LB and spotted around 5 µl of diluted cultures on LB agar plate and incubated at 37°C. For colony counting, approximately 10⁻⁶ dilution cells were spread on corresponding selection LB agar plate. Viable colonies were counted and expressed as cfu/ml of cell counts. For growth curve analysis, cells were grown in LB medium containing respective stressors and OD₆₀₀ readings were followed at every hour till the cultures reached stationary phase and graph was plotted with OD₆₀₀ versus time.

DNA constructs

cDNA of eukaryotic PABPN1 was obtained from Origene (PCMV6-AC with cloned PABPN1). Full-length PABPN1 was PCR amplified and cloned at the EcoRI and HindIII sites of pBAD18 plasmid to generate pBAD18-PABP, and at the Ndel and HindIII sites of pET28a vector to generate pET-PABP. PA-tail binding mutations (Y175A) were introduced by site directed mutagenesis as described earlier (Sudheesh et al., 2019). hfq and cspE were PCR amplified from E. coli chromosomal DNA and cloned in EcoRI and Xhol sites of pET28a (pET-hfq and pET-cspE) vector respectively for recombinant protein purification. hfq and cspE were also cloned at the EcoRI and HindIII sites of pDAB18 plasmid to generate pBAD18-hfq and pBAD18-cspE. List of primers used for PCR amplification, site directed mutagenesis, inverse PCR and cloning are shown in Table S5.

Protein purification

Recombinant proteins were expressed using pET28 plasmid constructs, overexpressed in BL21(DE3) by inducing with 1 mM isopropyl thiо-ß-D-galactoside at 18°C. Cells were lysed in ice-cold lysis buffer (20 mM Tris and 200 mM NaCl) and purified by Ni-NTA affinity chromatography as described previously (Sudheesh et al., 2019). The purified proteins were dialyzed in protein storage buffer (20mM Tris-HCl and 100 mM NaCl), concentrated using polyethylene glycol (PEG 20000 mw), snap frozen and stored in -80°C.
Generation of chromosomal insertion

Chromosomal insertion of PABPN1 in the receiver strains MG1655 or pcnB mutant (MG1655ΔpcnB759::kan) strains was carried out using λ-Inch technology as described earlier. We used the MG1655 bearing pBAD18-PABP as donor strain and λ phage integration resulted in the insertion of a region from near ori till bla gene encompassing the cloned PAB from the pBAD18-PABP. Integration occurred in the intergenic region between the gal and bio operons that flanks the inserted PABP-bla gene in the chromosome of receiver strains without disruption of any functional genes generating MG-PABP or pcnB-PABP strains respectively.

Antibiotic sensitivity assay and plasmid copy number analysis

Antibiotic sensitivity assay was carried out with wild-type MG1655, and pcnB mutant strains transformed with pBAD18 or pBAD18-PABP plasmids, and streaked on LB-agar plates in the presence of increasing ampicillin concentration (100 µg/ml to 2000 µg/ml). Alternatively, stable chromosomally inserted strains were transformed with pBAD18 plasmid. To analyze the plasmid content and copy number, above strains were transformed with another an RNAI independent non-colE1 plasmid pCL1920. From the cultures collected at different OD_{600} in LB medium, plasmids were isolated by alkaline lysis, digested both pBAD18 and pCL1920 plasmids with NdeI, and analyzed on a 1% agarose gel. Copy numbers was calculated by quantifying the amount of plasmids as described earlier.

Immunoblotting

For Immunoblotting experiments, cell lysates or IP eluates were resolved on a SDS-PAGE and blotted using specific antibodies on a PVDF membrane as described earlier. In IP experiments, input was loaded at an amount that is equivalent of 10% of the IP sample.

RNA immunoprecipitation

RIP experiments were carried out after cross-linking total cellular proteins with nucleic acids using 1% formaldehyde in MG1655 and MG-PABP cells as described earlier. The immunoprecipitated samples were eluted and digested with DNAseI, and associated RNA was then detected using one step RT-PCR reaction with gene specific primers. List of primers used for RIP analysis are shown in Table S5.

Immunofluorescence and imaging

Immunostaining was carried out as describe earlier. MG1655, MG-pcnB, and MG-PABP cells were grown upto 0.5 OD_{600} and treated with NaCl and MNNG for one hour. Cells were then harvested at 6000 rpm for 5 minute at 4°C, fixed in ice-cold 80% methanol and 16% paraformaldehyde, followed by permeabilisation with 10 µg/ml of lysozyme. Cells were then mounted on cover slips, pre-treated with 0.1% of poly-L-lysine, blocked with 2% BSA and stained with DAPI. Slides were then imaged on a Nikon A1R Laser scanning confocal microscope as described earlier. Quantification of the surface area from the phase contrast images was carried out using NIS elements software.

Crystal violet assay for biofilm formation

Cells were grown in 96-well microtiter dishes till OD_{490} = 1. Wells were washed with double distilled water after carefully removing the un-adhered cells. Cells remaining in the wells were then stained with 0.1% crystal violet solution in water for 20 minutes as described previously. Unbound dye was then washed off with autoclaved water and the dye bound biofilms were solubilized in 100% DMSO for 20 minutes, and absorbance was measured at 595 nm.

Biotinylation pull-down

For biotinylation pull down, cell lysates were prepared from ~10 ml culture of MG1655 or MG-PABP and cells at OD_{600} of ~1.00 were lysed in lysis buffer (10 mM HEPES, pH-7, 200 mM NaCl, 1% Triton-X 100, 10 mM MgCl₂, 1 mM DTT and protease inhibitor) as described earlier. Briefly, ~30 µg of biotinylated RNA was incubated with 60 µl of pre-washed avidin-agarose beads at 4°C for 1 hr. Agarose beads were washed three times with RNA binding buffer (10 mM Tris HCl, 0.1 M KCl and 10 mM MgCl₂), incubated with 5 mg total protein equivalent cell lysates overnight at 4°C. Agarose-beads were collected and washed and proteins were eluted using SDS loading buffer after boiling for 10 minutes at 95°C. Associated proteins were analyzed by Western blotting using specific antibodies.
Quantitative real time PCR (qRT-PCR) and half-life measurement

Bacterial cells were grown in 37°C, harvested at OD600 of ~0.6 and total RNA was isolated by trizol method as described earlier (Chomczynski, 1993) and quantified by A260 measurement. 2.5 μg of total RNA was used to synthesize first strand cDNA with random hexamers and MMLV reverse transcriptase (Invitrogen). qRT-PCR was performed with gene specific primers and quantified with CFX96 multi-color system using SYBR Green Supermix (Bio-Rad) as described previously (Sudheesh et al., 2019). Single-product amplification was confirmed by melt-curve analysis and primer efficiency was near 100% in all experiments. Quantification is expressed in arbitrary units and target mRNA abundance was normalized to the expression of dxs or rrsA with the Pfaffle method as described earlier (Sudheesh et al., 2019).

For the half-life measurement of mRNA, cells were treated with 500 μg/ml rifampicin to inhibit transcription and harvested at multiple time points (0 to 20 min) post-rifampicin treatment as described earlier (Sudheesh et al., 2019). RNA was isolated from each cells collected from each time points (0, 1, 2, 4, 6, 8, 12, 16, 24 min). qRT-PCR was carried out and half-life (T1/2) was measured as described earlier by following the decrease in % mRNA level over time with 0 time point taken as 100% of each gene expression (Sudheesh et al., 2019). List of primers used for qRT-PCR analysis are shown in Table S5.

3’-RACE assay

Total RNA was isolated from MG1655, MG-pcnB and MG-PABP using trizol reagent. For 3’-RACE assay, first strand CDNA was synthesized using MMLV reverse transcriptase and 2-μg of total RNA with Adapter primer at the 5’-end with as described previously (Sudheesh et al., 2019). cDNA was further amplified using genes specific forward primer and AUAP reverse primer. Sequence of Adapter primer, AUAP primer, and gene specific primers are shown in Table S5.

In vitro polyadenylation assay

In-vitro polyadenylation assay was carried out using recombinant His-PAPI on universal A5 (UAGGGA)16(A15)(Mellman et al., 2008) template with radiolabelled [α-32P]-ATP in a PAP assay buffer (250 mM NaCl, 50 mM Tris-HCl, 10 mM MgCl2, pH 7.9 at 37°C) as described earlier (Mellman et al., 2008). RNA products were analyzed on a 6% urea denaturing poly-acrylamide gel and visualized by phosphor imaging.

In vitro RNA synthesis, EMSA and nuclease protection assay

PA-tailed (RNAI-A16) and non(A)-tailed (RNAI-A0) transcripts were prepared by in vitro-transcription using PCR products amplified with a forward primer containing T7 promoter sequence. In the reverse primer 16 Ts were incorporated to generate a short PA-tail of 16 nucleotides whereas for A0, no overhangs were attached. To prepare the biotin labeled RNAI, in vitro transcription was carried out with the same PCR product in the presence of biotin16-UTP to obtain a body labeled transcript as described earlier (Marin-Belajar and Huarte, 2013). EMSA experiments were carried out as described earlier in a 20 μl EMSA-binding buffer (10 mM Tris-HCl, pH-7.5, 1 mM EDTA, 50 mM NaCl, 0.5 mM MgCl2, 1 mM DTT) in the presence of 1 μg/ml bovine serum albumin, 50% glycerol with 0.1 nM radiolabeled RNA templates with increasing PABPN1, Hfq and CspE (3 to 30 nM) at room temperature for 30 minutes. The RNA-Protein complex was analyzed on a native polyacrylamide gel and visualized by phosphor imaging.

For nuclease protection assay, active cellular extracts were prepared in a cell lysis buffer containing 50 mM TrisHCl and 10 mM β-mercaptoethanol and digestion of substrate RNAI-A16 or RNAI-A0 was carried out as described earlier (Feng et al., 2001). ~5 nM of radiolabelled in vitro transcribed RNA substrates were incubated with 4 μg equivalent of each active cell lysates for various time points as indicated. The residual RNA was then purified by phenol-chloroform, precipitated and analyzed on a urea-denaturing acrylamide gel and visualized by phosphor imaging.

Genome wide RNA-Seq analysis

RNA extracted from pellets of MG1655, MG-PABP, MG-pcnB were analyzed for QC using Bioanalyser and Qubit. Library preparation and deep sequencing of each sample was performed at commercially available genomics facility at the Genotypic Technology, Bangalore-India (https://www.genotypic.co.in). 4 μg of QC
passed total RNA was used for ribo-depletion using RiboMinus Bacterial Kit (Invitrogen). Further, 100 ng of Qubit quantified ribo-depleted RNA was taken for Transcriptome library preparation according to SureSelect Strand-Specific RNA Library Prep Kit protocol outlined in “SureSelect Strand-Specific RNA Library Prep for Illumina Multiplexed Sequencing” (Illumina). Briefly, the RNA was fragmented for 4 minutes at 94°C in the presence of divalent cations and first strand cDNA was synthesized. The single stranded cDNA was cleaned up using High Prep (Magbio, Cat # AC-60050). Strand specificity was maintained by the addition of actinomycin D. Second strand cDNA was synthesized and end repaired using Second Strand Synthesis using End Repair mix. The cDNA was cleaned up using High Prep (Magbio, Cat # AC-60050). Adapters were ligated to the cDNA molecules after addition of “A” base. High Prep cleanup was performed post ligation. The library was indexed and enriched for adapter ligated fragments using 10 cycles of PCR. The prepared library was quantified using Qubit and validated for quality by running an aliquot on High Sensitivity Bioanalyzer Chip (Agilent). 150 (75X 2) bp Paired End Sequencing was carried out on Illumina platform to generate 20-25 million PE Reads per sample. Bioinformatics analysis was performed with algorithms for Alignment Statistics, Reference alignment with depth statistics, digital gene expression, and DEseq plot read counts. The raw data generated was checked for the quality with FastQC and pre-processed to remove adapter sequences and low quality bases. It was then aligned using Tophat-2.0.133 (Trapnell et al., 2009) to E. coli (K-12) (Ensembl https://bacteria.ensembl.org/Escherichia_coli_str_k_12_substr_mg1655_gca_000005845/Info/Index) reference genome. Transcript assembly was done using Cufflinks-2.2.1 (Trapnell et al., 2010) which assembles transcripts, estimates their abundances, and tests for differential expression and regulation in RNA-Seq samples. Then, using cuffmerge, cufflinks assemblies were combined followed by differential gene expression analysis using “Cuffdiff” to obtain the significant changes in transcript expression (Trapnell et al., 2010). Uniprot knowledge base was used to annotate the genes for Gene Ontology (GO) (Boutet et al., 2012). Differential gene expression studies were carried out between PABP N1 expressed group versus wild type MG1655 samples. Differentially expressed genes were further annotated for the protein name and Gene Ontology and heat map generated as a representation of the expression values (as colors). The raw RNA-Seq data of our study has been deposited in the NCBI GEO sequencing data repository.

**PolyA+ RNA isolation and PA-tail assay screening**

Bacterial cells were grown in 37°C, harvested at OD600 of ~0.6 and total cellular PA-plus RNA was isolated using Oligotex PA-plus RNA isolation kit (Qiagen) as per the manufacturers instruction. Eluted PA-plus RNA was quantified and employed for G-I tailing and cDNA synthesis for downstream 3'-end PA-tail assay (PA-tail assay kit, Affymetrix) as described previously (Kusov et al., 2001; Patil et al., 2014). For G-I addition at the 3'-end, ~500 ng of purified PA-plus RNA was incubated with yeast PAP in the presence of guanosine triphosphate (GTP) and inosine triphosphates (ITP) at 37°C for one hour. First strand cDNA was then synthesized from G-I tailed RNA using G-I tail specific reverse primer. cDNA was then used for both qualitative and quantitative PCR. For PCR amplification a gene specific forward primer (GSP) and a universal reverse primer (URP) specific for the added G-I tail was used. Qualitative PCR was analyzed by agarose gel electrophoresis and quantitative analysis was carried out by real-time qPCR using similar primers and expressed as relative RNA levels. Schematic depicting the PA-tail assay system is shown in Figure S2F. Screening of cellular ~86 tRNAs and ~85 sRNAs was carried out by PA-tail assay using forward primers specific to each tRNA or sRNA and a universal reverse primer specific to the G-I tail addition as described above. Changes in the polyadenylated RNA levels were analyzed qualitatively by PA-tail assay coupled RT-PCR and quantitatively by real-time qPCR.

**Phylogenetic analysis**

We carried out an exhaustive literature analysis for the occurrence of poly(A) binding proteins, poly(A) polymerase and mRNA stabilization by polyadenylation across various branches of the evolutionary tree. Published literatures, NCBI protein search and protein databases such as pfam were used as reference to mark presence or absence of identified PAPs or PABP protein in organisms across different phyla in the evolutionary tree. We however ignored the hypothetical proteins and un-annotated proteins from the finding. Further, phylogenetic tree was later constructed with the available occurrence with representations from all the domains across all kingdoms.
QUANTIFICATION AND STATISTICAL ANALYSIS

All data were obtained from at least three independent experiments and are represented as mean ± standard error mean, SEM. The statistical significance of the differences in the mean is calculated using ANOVA with statistical significance at a p-value of less than 0.05. All Western blots show representative of at least from three independent blotting experiments. Quantifications of blots were carried out with Image J software.