A Shuttle Vector Based Transformation System for

*Pyrococcus furiosus*

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**ABSTRACT**

*Pyrococcus furiosus* is a model organism for analyses of molecular biology and biochemistry of archaea but so far no useful genetic tools for this species have been described. We report here a genetic transformation system for *P. furiosus* based on the shuttle vector system pYS2 from *Pyrococcus abyssi*. In the redesigned vector, the
pyrE gene from *Sulfolobus* was replaced as selectable marker by the 3-hydroxy-3-methylglutaryl coenzyme A reductase gene (HMG-CoA) conferring resistance of transformants to the antibiotic simvastatin. Use of this modified plasmid resulted in the overexpression of the HMG-CoA reductase in *P. furiosus*, allowing the selection of strains by growth in the presence of simvastatin. The modified shuttle vector replicated in *P. furiosus*, but the copy number was only one to two per chromosome. This system was used for overexpression of His6 tagged subunit D of the RNA polymerase (RNAP) in *Pyrococcus* cells. Functional RNAP was purified from transformed cells in two steps by Ni-NTA and gel filtration chromatography. Our data provide evidence that expression of transformed genes can be controlled from a regulated gluconeogenetic promoter.

**INTRODUCTION**

Several reports addressed the initial establishment of genetic techniques for the *Thermococcales*, a major group of hyperthermophilic archaea including the genera *Thermococcus* and *Pyrococcus*. The first experiments described used the plasmid pGT5 from *Pyrococcus abyssi*. This plasmid is only 3440 bp in size and replicates via a rolling circle mechanism (7). The archaeal plasmid was fused with a pUC19 vector to create a potential shuttle vector between *Escherichia coli* and *Pyrococcus furiosus* (1). This construct could be transformed in both organisms by CaCl2 treatment. Later, this construct was modified by introducing the alcohol dehydrogenase gene from *Sulfolobus solfataricus* as a selectable marker (3). The resulting plasmids pAG1 and
pAG2 were maintained for several generations in \textit{E. coli}, in the euryarchaeote \textit{P. furiosus} and also in the crenarchaeote \textit{S. acidocaldarius}. The presence of these plasmids in the two archaea conferred resistance to butanol and benzyl alcohol.

As the attempts to use this selection system for \textit{P. abyssi} failed, a new shuttle vector, pYS2, was created (17). This construct is also based on the archaeal pGT5 plasmid and a bacterial vector, pLitmus38. It contains the \textit{pyrE} gene of \textit{S. acidocaldarius}, a key enzyme of the pyrimidine biosynthetic pathway, as a selectable marker. For the transformation procedure a \textit{Pyrococcus} strain was used containing a \textit{pyrE} mutation which led to a uracil-auxotrophic phenotype. Using the shuttle vector pYS2 in combination with a polyethylene glycol-spheroplast method, it was possible to transform the \textit{pyrE} mutant of \textit{P. abyssi} to uracil prototrophy. Although the transformation frequency was very low, the shuttle vector was stably maintained at high copy number under selective conditions in both \textit{E. coli} and \textit{P. abyssi} (17).

A major breakthrough in the establishment of genetic tools for hyperthermophilic euryarchaeota was the development of a targeted gene disruption system by homologous recombination in \textit{Thermococcus kodakaraensis} KOD1 (23). A uracil-auxotrophic strain was converted with a disruption vector harboring the \textit{pyrF} marker within the \textit{trpE} gene to a uracil-prototrophic and a tryptophan-auxotrophic strain by double-crossover recombination. Due to the natural competence for DNA uptake, the high transformation efficiency and the high incorporation rate of DNA into its genome by homologous recombination, the system led to the identification of novel biochemical pathways, discovery of new enzyme functions and further
elucidation of proteins involved in the basic process of transcription (4, 11, 20, 22). A further improvement of this genetic system was the discovery that overexpression of the 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase gene is connected with the resistance against the antibiotic simvastatin (18). This selection system was first described in halophiles (15) and has the great advantage that there is no need for a certain host strain with a particular defect or auxotrophy toward an amino acid (18).

These new findings with respect to antibiotic resistance and the fact that the published shuttle vectors from *Pyrococcus* were never used in further investigations prompted us to redesign the pYS2 vector (17). In this article we describe the construction of a modified shuttle vector which allows overexpression of the HMG-CoA reductase in transformed *Pyrococcus* cells and leads to stable simvastatin resistant cells. Furthermore, we also demonstrate overexpression of subunit D of the RNA polymerase (RNAP) by introducing an additional copy of subunit D with a C-terminal His tag under the control of a regulated promoter.

**MATERIALS AND METHODS**

**Strains and growth conditions.** *P. furiosus* was cultivated under anaerobic conditions at 85°C in nutrient-rich medium based on ½ SME-medium and supplemented with different organic substrates (8). ½ SME-starch medium contained 0.1% each starch, yeast extract and peptone. For ½ SME-pyruvate medium, the starch was replaced with 40 mM Na-pyruvate. Gelrite (1%) was added for solidification of medium. The
antibiotic simvastatin (Toronto Research Inc., Toronto, Canada) was dissolved in ethanol and sterilized by filtration.

**General DNA manipulation.** *Escherichia coli* strain DH5α, used for vector construction and propagation, was cultivated at 37°C in Luria-Bertani (LB) medium. When needed, 100 μg/ml ampicillin was added to media. The vector pYS2 was provided by Prof. Gaël Erauso (Université de la Méditerranée, Marseille, France). Restriction and modification enzymes were purchased from NEB (Ipswich, USA). Plasmid DNA und DNA fragments from agarose gels were isolated using a plasmid mini or gel extraction kit from Qiagen (Hilden, Germany). Phusion High-Fidelity DNA polymerase from Finnzymes (Keilaranta, Finland) was used as a polymerase for PCR. DNA sequencing was performed by Geneart (Regensburg, Germany). Genomic DNA from *P. furiosus* wild-type and transformed strains was isolated using a DNeasy Blood & Tissue kit from Qiagen.

**Construction of the shuttle vectors pYS3 and pYS4.** The overexpression cassette for the HMG-CoA reductase gene from *P. furiosus* was constructed by replacing the native promoter with the strong promoter region (-250 to -1) of the glutamate dehydrogenase gene (*gdh*; (10)). The fusion of this promoter to the coding region of the HMG-CoA reductase (Primers: PF1848F 5’-ATGGAAATAGAGGAGATTATAGAG-3’ and PF1848BamHI 5’-ATCATCGGATCCTCATCTCCCAAGCATTTTATGAGC-3’) was done by PCR with overhanging ends at the reverse primer for the *gdh* promoter region *gdh*PromR-PF1848 (5’-CTCCTCTATTTCCATGTTCATCCCTCCAAATTAGGTG
As forward primer for the amplification \textit{gdhPromFBamHI} 
(5`-GGAACCGGATCCTTGAAAATGGAGTGAGCTGAG-3`) was used. The 
cassette was inserted into the pYS2 vector by replacing the \textit{BamHI} fragment 
containing \textit{pyrE} by the \textit{hmg-Co} reductase gene. The created vector pYS3 was 
sequenced and used later for transformation and further modification.

To obtain shuttle vector pYS4, RNA polymerase subunit D \textit{(rpoD)} was linked to the 
the fructose-1, 6 bisphosphatase \textit{(fbp)} promoter and inserted into pYS3. A His\textsubscript{6} tag 
was attached at the C-terminus of subunit D in addition. The promoter sequence of the 
\textit{fbp} was amplified from genomic DNA using the primers \textit{EcoRV-PF0613Pr-F} 
(5`-CTATTAGATATCTCCTTAACATTTCTCCAAA-3`) and \textit{PF0613Prom-R} 
(5`-CTGAACTTCAATTCCGGCCATTTTTTCACCTCCAGAAT-3`). Via the 
\textit{PF0613Prom-R} primer the promoter sequence had a 3` overhang for the fusion with 
the coding region of \textit{rpoD}. For the amplification of subunit D the primers \textit{PF1647-F} 
(5`-AAATGGCCGGAATTGAAGTTCAGATTCTTGA-3`) and \textit{PF1647-His-R} 
(5`-GTGATGGTGATGGTGATGAGAGGTCAATTTTTGAAGTTCAC-3`) were 
used. This step introduced the incorporation of the sequence for the His\textsubscript{6} tag at the 
C-terminus of \textit{rpoD}. The \textit{rpoD-His\textsubscript{6}} was fused with the terminating region of the 
\textit{histone A1} gene of \textit{P. furiosus}. (24). The primer pair \textit{His-PF1831Term-F} 
(5`-CATCACCATCACCATCACTGAAATCTTTTTTAGCACTT-3`) and 
\textit{PF1831T-EcoRV-R} (5`-TCAATTGATATCACCCTAGAAAAAGATAAGC-3`) 
created the terminating region of the \textit{histone A1} gene with a part overlapping the 
\textit{rpoD-His\textsubscript{6}} at the 5`-end that was used to fuse the fragments by PCR. Finally, the
construct was integrated into the pYS3 vector next to the hmg-CoA reductase cassette using the flanking EcoRV sites. The construction of the plasmid was verified by DNA sequencing.

Transformation of *P. furiosus*. *P. furiosus* cultures grown at 75°C to a cell density between 0.8 - 1.0 x 10⁸ per ml were used for transformation. For a transformation reaction the cells of 3 ml grown culture were collected anaerobically by centrifugation (10 minutes at 6000 g) and resuspended in a total volume of 100 µl transformation solution containing ½ SME (without KH₂PO₄), 40 mM Na-pyruvate, 4.7 mM NH₄Cl and 80 mM CaCl₂. The pH was adjusted to 7.0 with HCl. Cells were incubated at 4°C for 90 minutes under anaerobic conditions. After 30 minutes 0.5 pmol pYS3 or pYS4 were added. After a heat shock at 80°C for 3 minutes, the cells were again incubated for 10 min at 4°C and then cultivated in ½ SME-starch liquid medium in the presence of 10 µM simvastatin at 85°C for 48 h. Later, the cells were plated on ½ SME medium with starch as substrate and containing 10 µM simvastatin. The plates were incubated at 85°C for 48 h.

Growth properties of *P. furiosus* and *P. furiosus* pYS3 and pYS4 transformants.

To analyze resistance toward simvastatin, pYS3 transformed cells were cultivated in ½ SME-starch medium supplemented with 1, 5, 10, or 20 µM simvastatin at 85°C. Wild-type *P. furiosus* cells were also cultivated in ½ SME-starch medium at 85°C, but without simvastatin. Cell densities were measured at appropriate intervals. Cell counts were analyzed with a Thoma counting chamber (0.02-mm depth; Marienfeld, Lauda-Königshofen, Germany) under a phase-contrast microscope. To determine the
expression of subunit D under glycolytic or gluconeogenetic conditions, *P. furiosus*
pYS4 cells were grown either in ½ SME-starch or ½ SME-pyruvate medium in the presence of 10 µM simvastatin at 85°C.

Detection of RpoD and RpoD-His6 by western blot analysis. For the preparation of cell extracts 10 g of *P. furiosus* wild-type or *P. furiosus* cells transformed with pYS4 were resuspended in 30 ml buffer (40 mM HEPES, 500 mM NaCl, 10 mM imidazole, 15% glycerol, pH 7.5), sonicated on ice and treated with glass beads using a FastPrep-24 (M. P. Biomedicals, Irvine, USA) for complete cell lysis. After centrifugation (100,000 g for 1 h at 4°C) the protein concentrations of the clarified supernatants were determined by Bradford assays. For quantification of the expression levels of RpoD or RpoD-His6, western blots were done as previously described using polyclonal antibodies raised against recombinant subunits A’´ or D from *P. furiosus* (9). The signals were visualized using a Cy5-labelled secondary anti-rabbit antibody from Thermoscientific (Waltham, USA) and a fluorescence image analyzer (FLA-5000, Fuji, Japan).

Purification of RpoD-His6 and RNAP-His6. The cell extracts prepared as described in the previous section were applied onto 1-ml Ni2+-charged HisTrap HP columns (GE Healthcare). Bound proteins were eluted in one step using an elution buffer containing 300 mM imidazole instead of 10 mM. To separate free RpoD-His6 from the fraction incorporated into the RNAP, the eluate was loaded onto a Superdex 200 gel filtration column (GE Healthcare) equilibrated with 40 mM HEPES, pH 7.3, 250 mM KCl, 2.5 mM MgCl2, 0.5 mM EDTA, 20% glycerol. Aliquots of the fractions were analyzed for
RNAP activity using a specific *in vitro* transcription assay (10) and SDS-PAGE analysis.

**Southern blot analysis.** Total genomic DNA was digested with *Eco*RV and the resulting restriction fragments were separated on a 1% agarose gel. After electrophoresis, the DNA was transferred to a nylon membrane (Roche Applied Science, Mannheim, Germany) by capillary blot. A part of the *rpoD* gene was amplified by PCR using the primer pair RpoD500-F (5’-CCAACATTTGCAGTTGATGAAG-3’) and RpoD500-R (5’-CTCTTCGAATCTCTTTGGATGTAG-3’). This segment was used as probe to detect the RNAP subunit D gene in genomic and in plasmid DNA. The labelled probe was generated by the random primed method using the NEBlot kit (NEB, Ipswich, USA) in the presence of digoxigenin-11-dUTP (Roche Applied Science, Mannheim, Germany). After hybridisation the signals were detected using anti-digoxigenin antibodies conjugated with alkaline phosphatase according to the instructions of the producer (Roche Applied Science, Mannheim, Germany).

**RESULTS**

**Transformation in *P. furiosus* with a redesigned shuttle vector of pYS2.** The selection mechanism of the shuttle vector pYS2 is based on a uracil auxotrophic strain of *P. abyssi* which has a mutation in the *pyrE* gene (17). The plasmid contains a wild-type copy of the *pyrE* gene of *S. acidocaldarius* and successful transformation complements the uracil auxotrophy. As our attempts to construct a uracil auxotrophic
strain of \textit{P. furiosus} were not successful (data not shown) we redesigned the vector pYS2. In the new construct pYS3 the \textit{pyrE} gene was substituted by the \textit{hmg-CoA} reductase gene and for an efficient expression this gene was fused with the strong \textit{gdh} promoter from \textit{P. furiosus} (Fig. 1).

As overexpression of the HMG-CoA reductase led to the resistance against simvastatin in \textit{T. kodokaraensis} (18) we also analyzed the effect of various concentrations of simvastatin on the growth of \textit{P. furiosus}. In ½ SME-pyruvate medium supplemented with 5, 10, or 20 \textmu M simvastatin, growth was inhibited for only one day, if the cells were incubated at 95°C. In contrast, incubation at 85°C with similar concentrations prevented growth for three days. This indicates that the stability of simvastatin is dramatically decreased at 95°C, but 85°C seems to be an appropriate temperature for selection of transformants. This reduced incubation temperature still allows growth of \textit{P. furiosus} in a reasonable time in liquid as well as in solidified medium.

In the first experiments, the new construct pYS3 was used to transform \textit{P. furiosus} according to the published CaCl$_2$ procedure for \textit{T. kodakaraensis} with some minor modifications (23): The heat shock was performed for 3 minutes at 80°C instead of 45 seconds at 85°C and cells were incubated in the cold at 4°C instead at 0°C. Transformants were selected by growing cells for 48 h in liquid medium in the presence of 10 \textmu M simvastatin. Growth was only observed when cells were transformed with plasmid pYS3 and not when cells were treated in control reactions with transformation solution not containing the plasmid. The transformation
efficiency in liquid medium was approximately $5 \times 10^2$ transformants per µg pYS3 plasmid DNA. For the isolation of single transformants cells grown in liquid cultures were plated on culture medium containing 10 µM simvastatin. The plating efficiency of the transformants in the presence of 10 µM simvastatin was ~15% (the plating efficiency of WT cells on media not containing the antibiotics was ~78%).

A few simvastatin resistant colonies were selected and further analyzed for the presence of the plasmid by PCR amplification. To provide evidence that the plasmid was stably replicated in *Pyrococcus* the plasmid was isolated again from *Pyrococcus* after several transfers (4-5 times) of cells in fresh culture medium. Using this isolated plasmid DNA it was possible to successfully re-transform *E. coli* (data not shown).

This clearly demonstrates that this redesigned shuttle vector including the plasmid pGT5 from *P. abyssi* was also stably replicated as an external DNA element in *P. furiosus*.

**Induced expression of subunit D of the RNAP.** As next step it was analyzed whether the shuttle vector could be converted into an expression vector which allows the expression of proteins under the control of a regulated promoter. Subunit D of the archaeal RNAP was used as a model protein and an additional copy of this gene was inserted into the shuttle vector under the control of the fructose-1-6 bisphosphatase (PF0613) promoter (Fig. 1, pYS4). To allow a simple and rapid purification of the protein a His$_6$ tag at the C-terminus was introduced and for efficient termination of transcription the terminator from the histone gene *hpyA1* was linked to the 3’-end of the gene (24). The PF0613 promoter is repressed under glycolytic and induced under
The new construct pYS4 was transformed into *Pyrococcus* and a single colony was transferred into liquid medium and first cultivated under glycolytic conditions in the presence of starch. Later, the same culture was transferred to gluconeogenic conditions using a medium containing pyruvate as energy source. In each case the expression of subunit D was analyzed in crude extracts and compared with the wild-type by a western blot assay using antibodies against RNAP subunit D (Fig. 2). Identical amounts of RNAP were applied to gels used for western blots as shown by immunostaining using the antibody raised against RNAP subunit A' . Analysis of the crude extracts of the wild-type strain revealed only one signal corresponding to subunit D (Fig. 2, lanes 7-9). In contrast, the crude extracts of the transformants grown with starch (lanes 1-3) or pyruvate (lanes 4-6) contained an additional polypeptide migrating slightly slower than wild-type subunit D. This signal corresponding to the additional copy of subunit D encoded on the plasmid differed in size due to the existence of the His$_6$ tag at the C-terminus. The additional signal found in transformants was rather weak in cells grown with starch and much stronger in cells grown with pyruvate (Fig. 2, compare lanes 1-3 with 4-6). This clearly demonstrates that the promoter of the additional subunit D copy on the plasmid is strongly induced under gluconeogenic conditions and therefore this system is useful for regulated expression of proteins in *P. furiosus*.

Purification of archaeal RNAP by immobilized metal ion affinity
To analyze whether this modified subunit D containing a His$_6$ tag at the C-terminus also assembles into the archaeal RNAP, the crude extract of cells transformed with pYS4 was applied onto Ni-NTA columns. Specific bound proteins were eluted with a buffer containing 300 mM imidazole. To separate subunit D assembled into the archaeal RNAP from the free polypeptide the pooled subunit D-containing fractions from the Ni-NTA column were further purified by gel filtration chromatography. The RNAP containing fraction isolated by this two-step procedure from the transformant grown with pyruvate was compared with conventionally purified native RNAP (19) by gradient SDS PAGE and silver staining (Fig. 3A, lanes 2 and 3). Both RNAPs showed an almost identical pattern. This indicates that the overexpressed subunit D with the His$_6$ tag assembles into the RNAP and the whole enzyme can be isolated by immobilized metal ion affinity chromatography.

Starting with similar amounts of cells for purification, five-times more RNAP was isolated from transformant cells grown with pyruvate compared to cells grown with starch (Fig. 3A, compare lanes 4 and 5). As expected, RNAP was not enriched in similar fractions purified from extracts of the wild-type strain by the same procedure (Fig. 3A, lane 6). The band labelled with an asterisk was purified from wild-type cells as well as from the transformants after Ni-NTA affinity chromatography and Superdex 200 gel filtration. This multimeric polypeptide was not characterized in any detail here.

To check whether the affinity purified RNAP fractions (Fig. 3A, lanes 4 to 6) are functionally active these fractions were analyzed by \textit{in vitro} transcription experiments.
The affinity purified fractions were able to synthesize run-off RNA products from the 
*gdh* promoter in the presence of both archaeal transcription factors, TBP and TFB
((10); Fig. 3B, lanes 2 and 3). When one transcription factor or both were omitted, 
transcription was abolished (Fig. 3B, lanes 4 to 6). Taken together, these data indicate 
that subunit D with a C-terminal His\(_6\) tag assembles into the RNAP. As expected the 
amount of RNAP containing His6-tagged D that can be purified from a given amount 
of cells is higher when subunit D was overexpressed from the gluconeogenetic 
promoter. Furthermore, it is possible to specifically purify this fraction of the RNAP 
by Ni-NTA and size exclusion chromatography from the crude extract. The purified 
fraction is functionally active and not contaminated with TBP and TFB. As expected, 
this procedure did not allow the purification of RNAP without the C-terminal His\(_6\) tag 
at subunit D (Fig. 3B, lane 1).

**Copy number of pYS4 in *P. furiosus***. To determine the copy number of this shuttle 
vector in *P. furiosus* EcoRV-digested total DNA was analyzed by Southern blot 
experiments. The DNA sequence of subunit D was used as probe. When wild-type *P. 
furiosus* DNA was analyzed a 4.3 kb signal was identified. This exactly corresponds 
to the predicted size of a fragment containing subunit D in chromosomal DNA 
restricted with EcoRV (Fig. 4, lanes 2-6). When transformants were analyzed beside 
the genomic fragment an additional band with a size of 1.1 kb was observed (Fig. 4, 
lanes 7 to 11). As this signal was also present in the control lane with 
EcoRV-hydrolyzed plasmid DNA (lane 1) these results clearly demonstrate the
presence of plasmid pYS4 in transformed cells. The finding that the strength of both
signals arising from the genomic fragment and from the plasmid are in a similar range
indicates that the copy number of the plasmid is approximately in the same ratio as the
number of chromosomes of one cell. To exclude the possibility that the ratio between
plasmid and chromosomal DNA was changed during the DNA purification procedure,
the ratio between plasmid and chromosomal DNA was analyzed in crude extracts in
addition. This experiment confirmed the results that the copy number of the plasmid
pYS4 in \textit{P. furiosus} was between one and two (data not shown).

\section*{DISCUSSION}

In this paper we describe a shuttle expression vector system for \textit{P. furiosus} and \textit{E. coli}
allowing the regulated expression of proteins in \textit{Pyrococcus}. We have redesigned a
published shuttle vector of \textit{P. abyssi} using the overexpression of the HMG-CoA
reductase as a selection marker which confers resistance to the antibiotic simvastatin
as described earlier for \textit{T. kodakaraensis} (18). The copy number of the new vectors
pYS3 and pYS4 was dramatically reduced in comparison to the pYS2 shuttle vector
used in \textit{P. abyssi}. The copy number of the shuttle vector pYS2 was 20 to 30 copies per
chromosome and was therefore in the same range as described for the wild-type pGT5
plasmid from \textit{P. abyssi} (6, 17). At present we have no explanation for the dramatic
reduction of the copy number to one or two per chromosome in \textit{P. furiosus}. As in our
construct the transcription of the \textit{hmg-CoA} reductase gene occurs in opposite direction
to the replication of the plasmid, we have also analyzed whether insertion of the
hmg-CoA reductase gene in the opposite direction affects the copy number. First experiments indicate that the transcriptional orientation of the hmg-CoA reductase gene does not influence the copy number of the plasmid (data not shown). We assume that the maintenance of this plasmid in P. furiosus is mainly driven by the antibiotic resistance and the mechanism responsible to maintain a certain copy number in P. abyssi is absent in P. furiosus. A reduced copy number in a different host was also observed when plasmids pAG1 and pAG2 from P. abyssi were transferred to P. furiosus (1) or when a plasmid from Thermococcus nautilus was transferred to T. kodakaraensis (21).

Although the copy number of the shuttle vector described in this paper is low, the possibility to transform P. furiosus and to use this shuttle vector for the regulated expression of plasmid-encoded genes now allows the development of a genetic system. Our recent results suggest that it will be also possible to mutate the chromosome of P. furiosus using the overexpression of the HMG-CoA reductase as a selection marker against simvastatin (data not shown). But also the shuttle vector based regulated expression system offers novel intriguing possibilities for future developments. As we could successfully demonstrate that this system allows overexpression of a His\textsuperscript{6} tagged subunit D under the control of a gluconeogenetic promoter whose expression was dependent upon the substrate in the growth medium, the system described here can be used for production of recombinant proteins in P. furiosus. It could be an alternative system for expression of proteins which are difficult to produce in E. coli, especially for proteins of hyperthermophiles which might have a low propensity to
fold properly at low temperature like the Sor (sulfur oxygenase/reductase) protein from *Acidianus ambivalens* which was produced with higher efficiency in a hyperthermophilic *Sulfolobus* expression system than in *E. coli* (2).

We provide also evidence that this system can be used to isolate an active fraction of RNAP in a two step purification procedure when a His$_6$ tagged additional copy of subunit D was overexpressed in *Pyrococcus* cells. This system allows to overexpress mutant subunits in *Pyrococcus* and to isolate the RNAP containing mutations for structure function analyses as described for *T. kodakaraensis* (11, 21). Using a low level of expression of a particular subunit from the PF0613 promoter (growth on starch) it should be also possible to introduce point mutations into functional important regions of this subunit. Therefore, this system is a perfect complementation of our previously described system for the reconstitution of the 11-subunit RNAP from individual subunits *in vitro* (19). Furthermore, this system will be useful for the construction of a reporter gene assay, which should allow a rapid *in vivo* analysis of promoter sequences or of regulatory DNA elements in *P. furiosus*. Taken together, the presented shuttle vector based transformation system for *P. furiosus* is an important first step to establish a complete genetic toolbar for one of the hyperthermophilic key organisms in archaeal research for analysis of recombination (26), replication (5, 14), transcription (25) and metabolism (12).

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REFERENCES


FIGURE LEGENDS

FIG. 1. Schematic diagram of the shuttle vectors pYS2, pYS3 and pYS4. To create pYS3 the pyrE marker was substituted from pYS2 via BamHI restriction sites with a simvastatin resistance cassette. This enables the overexpression of the HMG-CoA reductase under the control of the gdh promoter. The plasmid pYS4 contains an additional fragment for the expression of a His6 tagged version of RNAP subunit D. This further copy is under the control of the fbp promoter and followed by the hpyA1 terminator sequence. The fragment was inserted into the EcoRV restriction site of pYS3.

FIG. 2. Comparison of the expression of wild-type and His6 tagged subunit D encoded on plasmid pYS4 by a western blot analysis. To analyse the relative amount of subunit D in the transformant grown with starch or pyruvate and in wild-type cells grown with starch we used comparable amounts of RNAP in each extract. For titration of equal amounts of RNAP we used an anti-subunit A’’ antibody (upper signal). The corresponding amounts of proteins in crude extracts from the transformant grown with starch were 218 ng, 436 ng and 872 ng (lanes 1-3). Lanes 4 to 6 contained the transformant grown with pyruvate with protein amounts of 1122 ng, 2244 ng and 4488 ng. Wild-type crude extracts were 166 ng, 332 ng and 664 ng protein (lanes 7-9). For identification of subunit D we used an anti-subunit D antibody (lower signals). Lane 10 contains a prestained size marker (Pager Ruler, Fermentas) and lane 11 50 ng of purified RNAP (with a His tag at subunit D).
FIG. 3. Purification and functional analysis of the RNAP. (A) Silver stained gradient SDS gel (4–20 %) of 2 µg purified RNAP from Pyrococcus. Lane 2 contains a wild-type RNAP, purified as described previously (10). Lane 3 contains the RNAP from the transformant grown with starch and purified with Ni-NTA and gel filtration chromatography. The corresponding fractions (identical volumes) of similar amounts of two-step purified cell extracts from wild-type (lane 4; without RNAP activity) and transformants grown with starch (lane 5) and pyruvate (lane 6) were analyzed on a 10 % SDS gel. The band labeled with an asterisk is an unknown protein which was copurified in all three extracts. The corresponding subunits of the RNAP are indicated, lane 1 contains a molecular weight marker. (B) In vitro transcription with the purified RNAP fractions. The RNAP fractions from the transformants grown with starch (lane 2) or pyruvate (lane 3) and the wild-type (lane 1) were used to transcribe the gdh promoter in the presence of the archaean transcription factors TBP and TFB. Lanes 4 to 6 contain control experiments without the transcription factors TBP or TFB or both. The transcription assays were performed as described previously (19).

FIG. 4. Analysis of the copy number of pYS4 in P. furiosus by a southern blot. Identical amounts from 25 ng to 400 ng of EcoRV-digested genomic DNA of P. furiosus wild-type (lanes 2-6) and pYS4 transformant (lanes 7-11) were used and analyzed with the rpoD probe. Lane 1 contained the EcoRV digested vector pYS4 as a positive control.