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A BROYDEN RANK $p + 1$ UPDATE CONTINUATION METHOD WITH SUBSPACE ITERATION∗

T. L. VAN NOORDEN†, S. M. VERDUYN LUNEL‡, AND A. BLIEK§

Abstract. In this paper we present an efficient branch-following procedure that can be used not only to compute branches of periodic solutions of periodically forced dynamical systems but also to determine the stability of the periodic solutions. The procedure combines Broyden’s method with a subspace iteration method to determine the dominant eigenvalues. The method has connections with the hybrid Newton–Picard methods developed by Lust et al. in [SIAM J. Sci. Comput., 19 (1998) pp. 1188–1209]. A convergence analysis of the procedure is presented. The method is applied to the computation of periodic states of a reverse flow reactor, and its performance is compared with two variants of the hybrid Newton–Picard method.

Key words. Broyden’s method, periodically forced partial differential equations, reverse flow reactor, continuation of periodic solutions

AMS subject classifications. 35B10, 35B32, 65M12, 65P30

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1. Introduction. This paper is devoted to the efficient computation of periodic solutions of periodically forced partial differential equations and the determination of their stability. The partial differential equations of interest model cyclically operated chemical systems. Examples of such dynamical systems are pressure swing separation [19], the reverse flow reactor [8], and the pressure swing reactor [9].

In order to investigate the behavior of periodic dynamical systems numerically, we first have to discretize the equations. For a shooting method, first the spatial variable of the equations is discretized. This results in a large system of $N$ time periodic ordinary differential equations. This system can abstractly be written as

$$\dot{x}(t) = f(t, x(t), \lambda), \quad x(t) \in \mathbb{R}^N, \quad \lambda \in \mathbb{R}, \quad \text{with} \quad f(t, \cdot, \cdot) = f(t + 1, \cdot, \cdot).$$

The assumption in (1.1) that the period length equals 1 is not a limitation. In fact the time variable $t$ can always be rescaled in order to assume that the period of the periodic forcing in $f$ is equal to 1.

The map $F : \mathbb{R}^N \times \mathbb{R} \to \mathbb{R}^N$ that for a given $\lambda$ assigns to the initial data at time zero, $x(0) = x_0$, the value of the solution after one period, i.e., the state $x(1)$, is called the Poincaré or period map of (1.1),

$$F(x_0, \lambda) = x(1, x_0),$$

where $x(t, x_0)$ denotes the solution of (1.1) with initial condition $x(0) = x_0$. So, evaluating the map $F$ is equivalent to integrating system (1.1) over one time unit starting with a given initial condition $x_0$.

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A 1-periodic state of the chemical system is a 1-periodic solution \( x(t) \), i.e., a solution of (1.1) with \( x(0) = x(1) \), and therefore the initial condition of a 1-periodic solution of (1.1) is a solution of the fixed point equation

\[
(1.3) \quad x = F(x, \lambda).
\]

In a similar way we can define 2-periodic, 3-periodic, etc., states of the chemical system.

We define the Jacobian of \( F \) at the point \( x = (x, \lambda) \) to be the square matrix \( J_F(x, \lambda) \) given by

\[
J_F(x, \lambda) = \partial F / \partial x(x, \lambda).
\]

The stability of a solution \( x^* \) of (1.3) is determined by the eigenvalues of the Jacobian \( J_F(x^*, \lambda) \) at the fixed point \( x^* \).

If we consider solutions \( x = (x, \lambda) \) of the equation \( F(x, \lambda) = x \), it is known that these solutions form in general, at least locally, a curve in \( \mathbb{R}^{N+1} \); see, for example, [6]. Pseudo-arc-length continuation [10] allows us to compute this curve also past turning points. Suppose we have a solution \( x_0 = (x_0, \lambda_0) \) of \( F(x, \lambda) = x \), as well as the normalized direction vector of the solution curve at \( (x_0, \lambda_0) \), which we will denote by \( \dot{x}_0 = (\dot{x}_0, \dot{\lambda}_0) \). Pseudo-arc-length continuation consists of solving the following equations for \( x_1 \) and \( \lambda_1 \):

\[
(1.4) \quad \begin{cases}
F(x_1, \lambda_1) - x_1 = 0, \\
\dot{x}_0^T(x_1 - x_0) + \dot{\lambda}_0(\lambda_1 - \lambda_0) - s = 0,
\end{cases}
\]

where \( s \) is the step size and \((x_1 - x_0)^T\) is the transpose of \((x_1 - x_0)\). Depending on how the equations (1.4) are solved, it is sometimes expensive to compute the direction vector \( \dot{x}_0 = (\dot{x}_0, \dot{\lambda}_0) \). In these cases a fair approximation can be obtained from

\[
(1.5) \quad \dot{x}_0 \approx \frac{(x_0 - x_{-1})}{|x_0 - x_{-1}|},
\]

where \( x_{-1} \) is a previously computed solution on the curve.

There exist various ways to obtain numerical solutions to (1.3) or (1.4). When a periodic state \( x(t, x^*) \) is stable, i.e., all the eigenvalues of the Jacobian \( J_F(x^*, \lambda) \) are inside the open unit disc \( D := \{ z \in \mathbb{C} : |z| < 1 \} \), then a solution to (1.3) can be obtained by integrating (1.1) on a long time interval, starting with an initial condition that is sufficiently close to the periodic solution. This approach is equivalent to a Picard iteration of the map \( F \). This method is, in many cases, not feasible, since the convergence to a periodic state of a dynamic simulation may be very slow [19]. Furthermore, only stable periodic states can be obtained by this approach.

A Newton-type method, applied directly to the fixed point equation (1.3), would allow us to compute both stable and unstable periodic states. However, such an approach involves the computation or approximation of the Jacobian \( J_F(x^*, \lambda) \) at various points \( x \), either by numerical differentiation or by solving the variational form of (1.1). Both approaches are very expensive when \( N \) is large [13, 3]. Note that although in many applications the Jacobian of \( f \) in the right-hand side of (1.1) is sparse, the Jacobian \( J_F(x, \lambda) \) does generally not have a sparse structure.

In numerical literature, a number of hybrid Newton–Picard methods have been developed; see [13, 7, 16]. These methods use the fact that for many physically interesting (infinite-dimensional) systems, the interesting dynamics actually occurs only in a low-dimensional subspace. Numerical evidence that this is also a feature for many periodic chemical processes is presented in [20]. A vector basis of this low-dimensional subspace is obtained from a subspace iteration algorithm. This approach
also yields eigenvalues that determine the stability. Furthermore, the action of the Jacobian \( J_F(x, \lambda) \) is computed only on the space spanned by this basis. It turns out that the theoretical convergence rate of the Newton–Picard method is linear.

In the chemical engineering literature other methods have been proposed to solve (1.3) or (1.4) more efficiently. It has been shown that Broyden’s method is very efficient in most cases [18, 8]. Broyden’s method uses only one evaluation of \( F \) per iteration and converges q-superlinearly. However, the disadvantage of Broyden’s method is that the approximations that the method produces to the Jacobian \( J_F(x^*, \lambda) \) of \( F \) at the periodic solution \((x^*, \lambda)\) are not accurate enough for the determination of the stability of the periodic solution.

In this paper we discuss a Broyden rank \( p + 1 \) update continuation method with subspace iteration (BSI rank \( p + 1 \) method, for short). The method combines the efficiency of Broyden’s method with the ability of the subspace iteration method to determine the stability of the computed solutions. The application of both the BSI rank \( p + 1 \) method and the Newton–Picard method to an example system shows that the BSI rank \( p + 1 \) method is approximately twice as efficient as the Newton–Picard method in terms of the number of evaluations of \( F \). The difference in efficiency lies mainly in the fact that the BSI method uses less evaluations of \( F \) per iteration than the Newton–Picard method. The example shows that the convergence rate and robustness of the two methods are comparable. We also consider the theoretical convergence properties of the Broyden rank \( p + 1 \) update method and show that an adapted version of the method is q-superlinearly convergent.

Since the value \( F(x_0, \lambda) \) is obtained by integrating a large system of differential equations over a period of time with initial condition \( x_0 \), the evaluation of \( F \) is a computationally expensive task. This means that for any method used to find a fixed point of \( F \), most of the computation time will be spent on the evaluation of \( F \). Thus the method that needs the least number of evaluations of \( F \) will be the most efficient method. Therefore the comparison between the BSI rank \( p + 1 \) method is made with respect to the number of evaluations of \( F \) needed to compute a branch of periodic solutions.

The organization of this paper is as follows. In section 2 we introduce the BSI rank \( p + 1 \) method and discuss the merits of the method in comparison with the Newton–Picard method. In section 3 we discuss the implementation of the BSI rank \( p + 1 \) method. In section 4 we analyze the theoretical convergence properties of the BSI rank \( p + 1 \) method. In section 5 we present the model equations for the cooled reverse flow reactor, and in section 6 we compute a branch of periodic states using the BSI rank \( p + 1 \) method, Broyden’s method, and the Newton–Picard method, and we compare the efficiency of all methods.

2. BSI rank \( p + 1 \) method. In the introduction we saw that, for the computation of a branch of periodic solutions of (1.1), we have to solve repetitively systems of the form

\[
\begin{align*}
F(x, \lambda) - x \\
w^T(x - y) + \kappa(\lambda - \mu) - s &= 0,
\end{align*}
\]

where \( w, y \in \mathbb{R}^N \) and \( \kappa, \mu, s \in \mathbb{R} \) are given. For convenience, we define the map \( G: \mathbb{R}^{N+1} \rightarrow \mathbb{R}^{N+1} \) by

\[
G(x) = G(x, \lambda) := \left( \frac{F(x, \lambda) - x}{w^T(x - y) + \kappa(\lambda - \mu) - s} \right),
\]
where \( x \) is the \((N + 1)\)-dimensional vector \((x, \lambda)\). An efficient way to compute solutions to the system \( G(x) = 0 \) is to use Broyden’s “good” method [1] (there also exist other variants of Broyden’s method; see, e.g., [5]). Broyden’s method produces approximations to a zero of \( G \) using the iteration scheme

\[
\mathbf{x}_{i+1} = \mathbf{x}_i - M_i^{-1} G(\mathbf{x}_i),
\]

with \( M_i \in \mathbb{R}^{(N+1)\times(N+1)} \) iterative approximations to \( \frac{\partial G}{\partial x}(\mathbf{x}_i) \) defined by

\[
M_{i+1} = M_i + \frac{(g_i - M_ip_i)p_i^T}{p_i^T p_i},
\]

where \( g_i = G(\mathbf{x}_{i+1}) - G(\mathbf{x}_i) \) and \( p_i = \mathbf{x}_{i+1} - \mathbf{x}_i \). This iteration scheme for \( M_i \) is derived from the following constraints on the update \( M_{i+1} \) of \( M_i \):

\[
M_{i+1}p_i = g_i,
\]
\[
M_{i+1}y = M_i y \quad \text{for all } y \perp p_i.
\]

Equation (2.5) can be viewed as a secant approximation of \( \frac{\partial G}{\partial x}(\mathbf{x}_{i+1}) \) in the direction \( p_i \). Note that the only information that is used in updating the approximation \( M_i \) at the new approximation \( \mathbf{x}_{i+1} \) is the function value \( G(\mathbf{x}_{i+1}) \). The method thus uses only one evaluation of \( F \) in each iteration. However, there is no guarantee that if the approximations \( \mathbf{x}_i \) converge to a solution \( \mathbf{x}^* \) of \( G(\mathbf{x}) = 0 \), the approximations \( M_i \) converge to \( \frac{\partial G}{\partial x}(\mathbf{x}^*) \). This means that the approximations \( M_i \) cannot be used to obtain good approximations of the eigenvalues of \( J_F(\mathbf{x}^*) \), which are needed for the determination of the stability of the solution \( \mathbf{x}^* \). In this section we discuss an approach to combine Broyden’s method with a subspace iteration algorithm [15] so that also approximations to the largest eigenvalues of \( J_F(\mathbf{x}^*) \) can be computed accurately.

For this approach we need not only the initial approximations \( \mathbf{x}_0 \) and \( M_0 \), required for Broyden’s method, but also initial approximations to the \( p \) largest eigenvalues of \( J_F(\mathbf{x}_0) \) together with an initial approximation \( V_0 \in \mathbb{R}^{N \times p} \) for the orthonormal basis of the subspace spanned by the \( p \) corresponding eigenvectors. In the first step of our new approach, we proceed as Broyden’s method and compute the next approximation to the solution of (2.1) with the same formula as before:

\[
\mathbf{x}_1 = \mathbf{x}_0 - M_0^{-1} G(\mathbf{x}_0).
\]

Now we would like to update \( M_0, V_0 \), and the approximations of the eigenvalues.

Let us first consider the updating of the eigenvalues and of \( V_0 \). We would like the updates of the eigenvalues to approximate the \( p \) largest eigenvalues of the Jacobian \( J_F(\mathbf{x}_1) \) in the new approximation \( \mathbf{x}_1 \) and the update \( V_1 \) to approximate the basis of the subspace spanned by the corresponding eigenvectors. Both the updates of the eigenvalues and the update of \( V_0 \) can be obtained by performing one (or more) subspace iteration(s). The version of the subspace iteration algorithm that we use is the version also used in [13]. The algorithm can be summarized by the following steps.

**Subspace iteration with projection.**

1. Set \( V_0 = V_0 \) and set \( j = 0 \).
2. Compute \( W_j = J_F(\mathbf{x}_1)\bar{V}_j \).
3. Compute \( \bar{S}_j = \bar{V}_j^T W_j \).
(4) Compute the Schur vectors $Y_j = [y_1, \ldots, y_p]$ of $S_j$; order them according to decreasing modulus of the corresponding eigenvalue.

(5) Compute $\tilde{V}_{j+1} = W_j Y_j$.

(6) Orthogonalize $\tilde{V}_{j+1}$.

(7) If $\tilde{V}_{j+1}$ is accurate enough, then stop and return $\tilde{V}_{j+1}$ and $W_j$. Else set $j = j + 1$ and go to step 2.

In this procedure the matrices $\tilde{V}_j$ will converge to an orthonormal basis of the subspace spanned by the eigenvectors and generalized eigenvectors associated with the $p$ largest eigenvalues of $J_F(x_1)$, ordered with respect to the modulus. The convergence properties of the subspace iteration algorithm are discussed in [15]. In fact the projection step 5 is not needed in the BSI rank $p + 1$ method, but in the present algorithm and implementation we have chosen to include the projection step in order for it to be compatible with the Newton–Picard method (see section 6).

In each iteration of the BSI rank $p + 1$ method, we perform only one iteration of the subspace iteration algorithm. This iteration provides us with new approximations to the $p$ largest eigenvalues of $J_F(x_1)$, with a new orthonormal basis $V_1 = \tilde{V}_1$, and with the matrix

\[
W_0 = J_F(x_1)\tilde{V}_0 = J_F(x_1)V_0.
\]

The matrix $W_0$ is very useful in updating $M_0$. If we define the matrices

\[
V := \begin{pmatrix} V_0 \\ 0 \end{pmatrix} \quad \text{and} \quad Z := \begin{pmatrix} W_0 - V_0 \\ W^TV_0 \end{pmatrix},
\]

then it is easily seen that

\[
\frac{\partial G}{\partial x}(x_1)V = Z.
\]

So, we can use (2.10), in the same way as the constraint (2.5) is used in Broyden’s method, to update the matrix $M_0$. Therefore, analogously to the Broyden rank one update, we would now like to update $M_0$ in such a way that

\[
M_1 V = Z,
\]

\[
M_1(x_1 - x_0) = (G(x_1) - G(x_0)),
\]

\[
M_1 q = M_0 q \quad \text{for all } q \text{ with } V^Tq = 0 \text{ and } (x_1 - x_0)^Tq = 0.
\]

As long as $(I - VV^T)(x_1 - x_0) \neq 0$, we have that the matrix $M_1$ characterized by (2.11)–(2.13) is well defined and unique. The matrix $M_1$ can be computed as

\[
M_1 = M_0 + \sum_{l=1}^{p} (z_l - M_0 v_l) v_l^T + (f - M_0 p)p^T
\]

\[
= M_0 + (Z - M_0 V)V^T + (f - M_0 p)p^T,
\]

where we have used the following notation:

\[
p = \frac{(I - VV^T)(x_1 - x_0)}{||(I - VV^T)(x_1 - x_0)||},
\]

\[
f = \frac{G(x_1) - G(x_0) - ZV^T(x_1 - x_0)}{||(I - VV^T)(x_1 - x_0)||}.
\]
It is easily verified that the matrix $M_1$ computed in (2.14) satisfies (2.11)–(2.13). We use the projected update $p$ and “projected” derivative vector $f$ instead of $x_1 - x_0$ and $G(x_1) - G(x_0)$, which are used in Broyden’s method, in order to obtain the orthogonal basis $v_1, \ldots, v_p, p$. It is this basis that is needed for the construction of $M_1$. Note that (2.11) and (2.12) are equivalent to (2.11) and $M_1 p = f$.

At this point we have computed the new updates $x_1, V_1, M_1$ and the new updates of the $p$ largest eigenvalues. By following the above procedure, starting with these updates, we can now obtain the next set of updates, and so on. This iteration of the above-described procedure characterizes the BSI rank $p + 1$ method. For convenience, we present three variants of this new procedure. The first one, which we call BSI1, can be summarized by the following steps.

BSI RANK $p + 1$ METHOD (BSI1).

1. Supply $M_0 \in \mathbb{R}^{(N+1) \times (N+1)}$, invertible; $V_0 \in \mathbb{R}^{N \times p}$, orthonormal; $x_0 = (x_0, \lambda_0) \in \mathbb{R}^{(N+1)}$, and set $i = 0$.
2. Solve $M_i s_i = -G(x_i)$ (computation of $G(x_i)$ costs one evaluation of $F$) and set $x_{i+1} = x_i + s_i$. Compute $S := J_F(x_{i+1}) V_i$ (costs $p$ evaluations of $F$).
3. Set $V := \begin{pmatrix} V_i \\ 0 \end{pmatrix}$ and $Z := \begin{pmatrix} S - V_i \\ w^T V_i \end{pmatrix}$.
4. Let $v_l$ and $z_l$ be the $l$th ($l = 1, \ldots, p$) column of, respectively, $V$ and $Z$. Set $v_{p+1} = \frac{(I-VV^T)s_{p+1}}{\| (I-VV^T)s_{p+1} \|}$ and $z_{p+1} = \frac{G(x_{p+1}) - G(x_i) - ZV^T S}{\| (I-VV^T)s_{p+1} \|}$. Compute $M_{i+1} = M_i + \sum_{l=1}^{p+1} (z_l - M_i v_l) v_l^T$.
5. Subspace iteration: Compute the ordered Schur-factorization $Y^T R Y$ of $V S$. Compute $V = SY$. Orthonormalize the columns of $V$ and put the result in $V_{i+1}$.
6. Set $i = i + 1$ and go to step 2.

This algorithm needs $p + 1$ evaluations of $F$ per iteration. In section 6 we will use the algorithm to compute a branch of periodic states of a reverse flow reactor.

Note that if for certain $i = i_0$, the update $s_i$ belongs to the subspace spanned by the columns of $V$, then the vectors $v_{p+1}$ and $z_{p+1}$ in step 4 of the algorithm above are not defined, and in that case we can update $M_i$ only with a rank $p$ matrix instead of a rank $p + 1$ matrix. More generally, if the angle between a certain update $s_i$ and subspace spanned by the columns of $V$ becomes small, we might encounter problems.

The sine of the angle between the update $s_i$ and the subspace spanned by the columns of $V$ is given by the quotient $\| (I - VV^T)s_i \|/\| s_i \|$. It turns out that this quotient is important in the convergence analysis of the BSI rank $p + 1$ method. To assure $q$-superlinear convergence of the method, this quotient has to be bounded away from zero. In the BSI rank $p + 1$ method as defined in algorithm BSI1, we cannot assure a priori that this quotient stays bounded away from zero. We can, however, adapt the BSI1 algorithm so that we can avoid problems. We next present two variants of the BSI rank $p + 1$ method, called BSI2 and BSI3, for which we can control the situation of a small angle.

In the first approach, BSI2, we use, in case the quotient $\| (I - VV^T)s_i \|/\| s_i \|$ becomes too small, an additional matrix-vector product in order to approximate the...
derivative of \( G \) in the direction \( ||(I - VV^T)s_i|| \), instead of using the secant update. In the second approach, BSI3, when the angle becomes too small, we use only a rank \( p - 1 \) update on a \((p - 1)\)-dimensional subspace of the \( p \)-dimensional subspace \( V \), i.e., the subspace spanned by the columns of \( V \). This \((p - 1)\)-dimensional subspace is chosen orthogonal to the update vector \( s_i \).

In both algorithms we introduce a parameter \( \kappa \), \( 0 < \kappa < 1 \), in order to control the influence of the sine of the angle between the update \( s_i \) and the subspace \( V \). In the limit \( \kappa = 0 \), the BSI2 and BSI3 algorithms reduce to the BSI1 method. Note that we only have to adapt step 4 in the BSI1 algorithm. Therefore we describe only this step for the BSI2 and BSI3 algorithms. The other steps remain unchanged.

BSI RANK \( p + 1 \) METHOD (BSI2).

(4) Let \( v_l \) and \( z_l \) be the \( l \)th \((l = 1, \ldots, p)\) column of, respectively, \( V \) and \( Z \).

Set \( v_{p+1} = (I - VV^T)x_i/s_i ||(I - VV^T)s_i|| \). Set \( c = ||(I - VV^T)s_i|| \). If \( 1/c < \kappa \), then compute

\[
\frac{\partial^2 G(x_{i+1})}{\partial x}v_{p+1} \text{, else set } z_{p+1} = \frac{G(x_{i+1}) - G(x_i) - ZV^T s_i}{||V^T s_i||}. \text{ Compute }
\]

\[
M_{i+1} = M_i + \sum_{l=1}^{p+1}(z_l - M_i v_l)v_l^T.
\]

Thus if the sine of the angle between the update \( s_i \) and the subspace \( V \) becomes smaller than \( \kappa \), we replace the “secant” update of \( M_i \) in the direction perpendicular to \( V \) by a directional derivative update. In this situation we need one extra directional derivative evaluation. The second alternative version of the BSI rank \( p + 1 \) method is given by the following step.

BSI RANK \( p + 1 \) METHOD (BSI3).

(4) Let \( v_l \) and \( z_l \) be the \( l \)th \((l = 1, \ldots, p)\) column of, respectively, \( V \) and \( Z \).

Set \( v_{p+1} = (I - VV^T)x_i/s_i ||(I - VV^T)s_i|| \). Set \( c = ||(I - VV^T)s_i|| \). If \( 1/c \geq \kappa \), then set \( z_{p+1} = \frac{G(x_{i+1}) - G(x_i) - ZV^T s_i}{||V^T s_i||} \) and compute

\[
M_{i+1} = M_i + \sum_{l=1}^{p+1}(z_l - M_i v_l)v_l^T.
\]

If \( 1/c < \kappa \), then set \( v_1 = V^T s_i/||V^T s_i|| \) and compute \( p - 1 \) vectors \( v_j \)

\((j = 2, \ldots, p)\) such that \((v_1, \ldots, v_p)\) forms an orthonormal basis for \( \mathbb{R}^p \). Now compute \( v_1 = V v_l \) for \( l = 2, \ldots, p \) and compute \( z_l = Z v_l \) for \( l = 2, \ldots, p \).

Set \( v_1 = s_i/||s_i|| \) and \( z_1 = \frac{G(x_{i+1}) - G(x_i)}{||s_i||} \), so that we can compute

\[
M_{i+1} = M_i + \sum_{l=1}^{p}(z_l - M_i v_l)v_l^T.
\]

In this case, if the sine of the angle between the update vector \( s_i \) and the subspace \( V \) becomes smaller than \( \kappa \), we update \( M_i \) with a rank \( p \) matrix, instead of the rank \( p + 1 \) matrix in the original method.

Note that if, in the BSI1 method, the angle between the update vector \( s_i \) and the subspace \( V \) stays bounded away from zero, then we can always set a value for \( \kappa \) such that all three variants, BSI1, BSI2 and BSI3, perform exactly the same steps. In section 6, we present results for each of the algorithms for various values of \( \kappa \).

In order to give an idea of the efficiency of the BSI rank \( p + 1 \) method, we compare its performance with the performance of two variants of a Newton–Picard method, the
continuation Newton–Picard (CNP) and continuation Newton–Picard Gauss–Seidel (CNPGS) methods, introduced in [13]. The CNP method needs $p+2l+4$ evaluations of $F$ per iteration, where $l$ is a parameter of the method that specifies the number of Picard iterations used in one full iteration of the Newton–Picard method. The CNPGS method needs $p+l+2$ evaluations of $F$ per iteration and so is less expensive than CNP. In [13] it is reported that CNP and CNPGS have the same asymptotic convergence rate, but that CNP happens to be more robust, i.e., has a larger domain of attraction, and thus allows larger step sizes in the continuation algorithm.

In [13], the subspace iteration is performed on $p+p_e$ (with $p_e = 2, 3, \text{or} 4$) vectors in order to accelerate the convergence of the subspace iteration procedure. In [13], also the number $p$ is varied during the computations in order to ensure that the subspace iteration is performed only on the subspace spanned by eigenvectors of which the corresponding eigenvalues are outside the region $D_\rho := \{ z \in \mathbb{C} : |z| < \rho \}$ (usually $\rho$ is taken to be 0.5). Here, in our setting, for both the Newton–Picard methods and the BSI method, we perform only the subspace iteration on $p$ vectors and keep this number fixed during the computations. Another difference between our approach and that of [13, 12] is that we do not use locking and deflation in the subspace iteration. Our approach with the dimension of $V$ kept fixed works very well for the cooled reverse flow reactor discussed in section 5 (see also [20]). However, for arbitrary dynamical systems, a procedure for varying $p$ as used in [16, 7] or in [13] is necessary.

In [13] it is reported that the Newton–Picard method converges linearly. Broyden’s method has theoretically better convergence properties. In [2] it is shown that Broyden’s method converges locally at least $q$-superlinearly, and in [5] it is shown that the method enjoys local 2N-step, $q$-quadratic convergence. In section 4 we will show that the BSI2 and BSI3 variants of the BSI rank $p+1$ method also converge $q$-superlinearly.

We also compare the BSI method with Broyden’s method followed by subspace iteration.

3. Implementation of the BSI rank $p + 1$ method. In this section we consider an efficient implementation of the BSI rank $p + 1$ method. In the previous section, we stated the BSI method in terms of approximations $M_i$ of the Jacobian of $G$. If we store the matrices $M_i$, then we need to solve the linear system $M_i x = -G(x_i)$ in each iteration of the BSI method. For the implementation of the BSI method it is therefore more convenient to store the inverses of the matrices $M_i$. Just as for Broyden’s method, we can easily update the inverse of $M_i$. In this way we do not need to solve a linear system in each iteration of the BSI method. The BSI1 method in terms of the matrices $H_i$ that are approximations of the inverse of the Jacobian of $G$ can be formulated as follows.

BSI rank $p + 1$ method (BSI1 reformulated).

1. Supply $H_0 \in \mathbb{R}^{(N+1) \times (N+1)}$, invertible; $V_0 \in \mathbb{R}^{N \times p}$, orthonormal; $x_0 = (x_0, \lambda_0) \in \mathbb{R}^{(N+1)}$, and set $i = 0$.

2. Compute

$$x_{i+1} = x_i + H_i G(x_i)$$

(costs one evaluation of $F$).

3. Subspace iteration: Compute $S := \frac{\partial F}{\partial x}(x_{i+1}) V_i$ (costs $p$ evaluations of $F$). Compute the ordered Schur-factorization $Y^T R Y$ of $VS$. Compute $V = SY$. Orthonormalize the columns of $V$ and put the result in $V_{i+1}$.
(4) Compute
\[
H_{i+1} = \begin{cases} 
B_p - \frac{(p + B_pf)p^TB_p}{p^TB_pf} & \text{if } p^TB_pf \neq 0 \\
\text{and } (I - ZZ^T)(G(x_{i+1}) - G(x_i)) \neq 0, \\
B_p & \text{otherwise},
\end{cases}
\]
where \(B_p\) is defined recursively by
\[
B_0 = H_i \
B_{l+1} = \begin{cases} 
B_l - \frac{(v_l + B_l(z_l)v_l^TB_l)}{v_l^TB_lz_l} & \text{if } v_l^TB_lz_l \neq 0, \\
B_l & \text{if } v_l^TB_lz_l = 0,
\end{cases}
\]
with \(v_l\) and \(z_l\) the \(l\)th column of, respectively,
\[
V := \begin{pmatrix} V_i \\ 0 \end{pmatrix} \quad \text{and} \quad Z := \left( \frac{\partial F}{\partial x}(x_{i+1}) - I \right) V_i,
\]
and where
\[
p = (I - VV^T)(x_{i+1} - x_i), \\
f = G(x_{i+1}) - G(x_i) - ZV^T(x_{i+1} - x_i).
\]
Set \(i = i + 1\) and go to step 2.

This algorithm is derived by using the Sherwood–Morrison formula recursively for the \(p + 1\) rank one updates of the matrices \(M_i\). It is easily checked that the inverse of matrix \(H_{i+1}\) computed in step 4 of the above algorithm satisfies (2.11)–(2.13). This shows that the two formulations of the BSI1 algorithm are indeed mathematically equivalent. Similarly, the BSI2 and BSI3 algorithms can also be stated in terms of approximations to the inverse of the Jacobian of \(G\).

An important issue for a numerical method is memory usage. The BSI rank \(p + 1\) method as discussed above needs to store the \(N \times N\) matrix \(H_i\). For the test problem presented in the paper, the value of \(N\) is not so large that the storage of \(H_i\) becomes a problem. For two-dimensional or three-dimensional model problems, however, the dimension of the discretized system \(N\) may become so large that it is impossible to store a full \(N \times N\) matrix. As an alternative to storing the full matrix \(H_i\), one can start with \(H_0 = I\), and then storing each rank \(p + 1\) update in the BSI rank \(p + 1\) method using \(2(p + 1)\) vectors. This approach, however, still leads to memory problems when the BSI rank \(p + 1\) method needs many iterations to converge.

To overcome this problem for Broyden’s method, there exist variants of the method with limited memory usage that in certain cases behave just like Broyden’s method [14]. The idea is to approximate \(H_i\), when the number of updates becomes too large, with a lower rank matrix so that the required storage remains limited. Such an approach might also be possible for the BSI rank \(p + 1\) method, but we do not pursue this matter further here. See [17] for a limited memory Broyden method that is also applied to computing periodic states of high-dimensional periodically forced dynamical systems.

The largest matrix that the Newton–Picard method requires to be stored is the \(N \times p\) matrix \(V_i\) that is an approximation to a basis of the subspace spanned by the eigenvectors corresponding to the largest \(p\) eigenvalues. Here the value of \(p\) is essentially independent of the discretization, so that the storage used by the Newton–Picard method depends only linearly on \(N\).
4. Convergence analysis of the BSI rank \( p + 1 \) method. For Broyden’s method, it is known that it converges \( q \)-superlinearly; see, for example, [4]. In this section we investigate whether this property also holds for the BSI rank \( p + 1 \) method. We will see that for the BSI1 method presented in section 2, this question cannot easily be answered. The two alternative variants BSI2 and BSI3, however, do enjoy \( q \)-superlinear convergence. Our numerical experiments in section 6 will show that the original BSI1 method behaves very much the same as the \( q \)-superlinearly convergent alternative variants.

For the convergence analysis of the BSI method, we follow the lines of the proof of the \( q \)-superlinear convergence of Broyden’s method as given in [4]. We assume that \( G : \mathbb{R}^{N+1} \rightarrow \mathbb{R}^{N+1} \) is continuously differentiable in an open convex set \( D \subset \mathbb{R}^{N+1} \).

We also assume that there exists \( x_0 \in \mathbb{R}^{N+1} \) and \( r, \beta > 0 \) such that

\[
B_r(x_0) := \{ \tilde{x} \in \mathbb{R}^{N+1} : \|\tilde{x} - x_0\| < r \} \subset D,
\]

\( G(x_0) = 0, \frac{\partial G}{\partial x}(x_0)^{-1} \) exists with \( \|\frac{\partial G}{\partial x}(x_0)^{-1}\| < \beta, \) and \( \frac{\partial G}{\partial x} \) is Lipschitz continuous on \( B_r(x_0) \) with Lipschitz constant \( \gamma \).

The first step is to prove that the generated sequence of matrices \( M_i \) is of bounded deterioration. This means that the approximation of the Jacobian can get worse, but slowly enough to ensure convergence to a zero of \( G \). This is the content of the following lemma.

**Lemma 4.1.** Let \( D \subset \mathbb{R}^{N+1} \) be an open convex set containing \( x_i \) and \( x_{i+1} \), with \( x_i \neq x_{i+1} \), where \( x_i \) is a solution of \( G(x) = 0 \). Let \( G \) be Lipschitz continuous on \( D \), and let \( x_0 \in D \). There exists a constant \( C \) such that

\[
\|M_{i+1} - J_*\| \leq \|M_i - J_*\| + C(\|x_{i+1} - x_*\| + \|x_i - x_*\|),
\]

where \( J_* \) denotes \( \frac{\partial G}{\partial x}(x_0) \), and where the matrices \( M_i \) are generated by the BSI2 or BSI3 algorithm.

**Proof.** From the description of the algorithms for BSI2 and BSI3, we have

\[
\|M_{i+1} - J_*\| = \left\| M_i - J_* + \sum_{l=1}^{p+1} (z_l - M_i v_l)v_l^T \right\|
= \left\| M_i - J_* + \sum_{l=1}^{p+1} (J_* v_l - M_i v_l)v_l^T + \sum_{l=1}^{p+1} (z_l - J_* v_l)v_l^T \right\|
= \left\| (M_i - J_*) \left( I - \sum_{l=1}^{p+1} v_l v_l^T \right) + \sum_{l=1}^{p+1} (z_l - J_* v_l)v_l^T \right\|
\leq \|M_i - J_*\| + \left\| \sum_{l=1}^{p+1} (z_l - J_* v_l)v_l^T \right\|.
\]

(4.1)

Here we have used the fact that, since all the vectors \( v_l \) are orthogonal and have norm equal to one, the matrix \( (I - \sum_{l=1}^{p+1} v_l v_l^T) \) is a Euclidean projection matrix and thus has norm one. The next step is to investigate the term \( \sum_{l=1}^{p+1} (z_l - J_* v_l)v_l^T \). First we split this term into two parts:

\[
\sum_{l=1}^{p+1} (z_l - J_* v_l)v_l^T = \sum_{l=1}^{p} (z_l - J_* v_l)v_l^T + (z_{p+1} - J_* v_{p+1})v_{p+1}^T.
\]
For the first part, we have

\[
\left\| \sum_{l=1}^{p} (z_l - J_* v_l) v_l^T \right\| = \left\| \sum_{l=1}^{p} \left( \frac{\partial G}{\partial x} (x_{i+1} - J_* v_l) \right) v_l^T \right\| \\
= \left\| \left( \frac{\partial G}{\partial x} (x_{i+1} - J_*) \right) \sum_{l=1}^{p} v_l v_l^T \right\| \\
\leq \left\| \frac{\partial G}{\partial x} (x_{i+1} - J_*) \right\| \leq \gamma ||x_{i+1} - x_*||.
\]

(4.2)

For the second part, we write

\[
|| (z_{p+1} - J_* v_{p+1}) || = \frac{||G(x_{i+1}) - G(x_i) - ZV^T s_i - J_*(I - VV^T)s_i||}{||(I - VV^T)s_i||} \\
= \frac{||G(x_{i+1}) - G(x_i) - J_* s_i - ZV^T s_i - J_* VV^T s_i||}{||(I - VV^T)s_i||} \\
\leq \frac{||G(x_{i+1}) - G(x_i) - J_* s_i||}{||(I - VV^T)s_i||} + \frac{||\frac{\partial G}{\partial x}(x_{i+1})VV^T s_i - J_* VV^T s_i||}{||(I - VV^T)s_i||} \\
\leq \gamma (||x_{i+1} - x_*|| + ||x_i - x_*||)||s_i|| + \gamma ||x_{i+1} - x_*||||s_i|| \\
2||(I - VV^T)s_i|| = \frac{\gamma}{\kappa} ||s_i||.
\]

If we have an upper bound for the quotient \( c = \frac{||s_i||}{||(I - VV^T)s_i||} \), we can also control this second part. For the BSI1 algorithm, we cannot assure that this quotient remains bounded, and therefore we cannot prove \( q \)-superlinear convergence for the BSI1 algorithm directly. For both the BSI2 and BSI3 algorithms, we observe that as long as

\[
1/c = \frac{||(I - VV^T)s_i||}{||s_i||} \geq \kappa,
\]

we have the following estimate:

(4.3) \( ||(z_{p+1} - J_* v_{p+1})|| \leq \frac{\gamma}{2\kappa} (||x_{i+1} - x_*|| + ||x_i - x_*||) + \frac{\gamma}{\kappa} ||x_{i+1} - x_*||.\)

If now \( 1/c = \frac{||(I - VV^T)s_i||}{||s_i||} < \kappa \), then for the BSI2 algorithm we have

(4.4) \( ||(z_{p+1} - J_* v_{p+1})|| = \left\| \frac{\partial G}{\partial x} (x_{i+1}) v_{p+1} - J_* v_{p+1} \right\| \leq \gamma ||x_{i+1} - x_*||,\)

where we have used the Lipschitz continuity of \( \frac{\partial G}{\partial x} \). Using (4.1), (4.2), (4.3), and (4.4), we obtain the following estimate:

\[
||M_{i+1} - J_*|| \leq ||M_i - J_*|| + \gamma (||x_{i+1} - x_*|| + \frac{\gamma}{2\kappa} (||x_{i+1} - x_*|| + ||x_i - x_*||)) \\
+ \frac{\gamma}{\kappa} ||x_{i+1} - x_*|| \\
\leq ||M_i - J_*|| + \gamma \left( \frac{3}{2\kappa} + 1 \right) (||x_{i+1} - x_*|| + ||x_i - x_*||).
\]

(4.5)
For the BSI3 algorithm we have instead of (4.4), the inequality

\[
|| (z_{p+1} - J_\ast v_{p+1}) || = \frac{|| G(x_{i+1}) - G(x_i) - J_\ast s_i ||}{|| s_i ||} \leq \frac{\gamma (|| x_{i+1} - x_\ast || + || x_i - x_\ast ||) || s_i ||}{2 || s_i ||},
\]

(4.6)

where we have used again the Lipschitz continuity of \( \frac{\partial G}{\partial x} \). Now using (4.1), (4.2), (4.3), and (4.6), we also arrive for the BSI3 variant of the BSI rank \( p + 1 \) method at the same estimate (4.5).

Now we are ready to prove the following theorem.

**Theorem 4.2.** Let \( G: \mathbb{R}^{N+1} \to \mathbb{R}^{N+1} \) be continuously differentiable in an open convex set \( D \subseteq \mathbb{R}^{N+1} \). Assume that there exists \( x_\ast \in \mathbb{R}^{N+1} \) and \( r, \beta > 0 \) such that \( B_r(x_\ast) \subseteq D \), \( G(x_\ast) = 0 \), \( \frac{\partial G}{\partial x}(x_\ast)^{-1} \) exists with \( || \frac{\partial G}{\partial x}(x_\ast)^{-1} || \leq \beta \), and \( \frac{\partial G}{\partial x} \) is Lipschitz continuous on \( B_r(x_\ast) \) with Lipschitz constant \( \gamma \). Then there exist positive constants \( \epsilon, \delta \) such that if \( || x_0 - x_\ast || \leq \epsilon \) and \( || M_0 - \frac{\partial G}{\partial x}(x_\ast) || \leq \delta \), then the sequence \( \{ x_i \} \) generated by the BSI2 or BSI3 algorithm is well defined and converges \( q \)-superlinearly to \( x_\ast \).

**Proof.** Lemma 4.1 allows us to invoke (part of) Theorem 8.2.2 in [4] to show that there exist positive constants \( \epsilon, \delta \) such that if \( || x_0 - x_\ast || \leq \epsilon \) and \( || M_0 - J(x_\ast) || \leq \delta \), then the sequence \( \{ x_i \} \) generated by the BSI2 or BSI3 algorithm is well defined and converges at least \( q \)-linearly to \( x_\ast \).

For the proof of the \( q \)-superlinear convergence, we can follow almost verbatim the appropriate part of the proof of Theorem 8.2.2 in [4] for the \( q \)-superlinear convergence of Broyden’s method. However, there is one minor modification that we will explain.

Define \( E_i = M_i - J_\ast \). It is not hard to see that (4.1) also holds for the Frobenius norm

\[
|| E_{i+1} ||_F \leq || E_i \left( I - \sum_{l=1}^{p+1} v_l v_l^T \right) ||_F + \left( \sum_{l=1}^{p+1} || z_l - J_\ast v_l ||_F v_l^T \right). 
\]

Because \( s_i \) is in the span of \( v_l, \ l = 1, \ldots , p + 1 \), and since we use the Frobenius norm, we also obtain the inequality

\[
|| E_{i+1} ||_F \leq || E_i \left( I - \sum_{l=1}^{p+1} \frac{s_l s_l^T}{s_l^T s_l} \right) ||_F + \left( \sum_{l=1}^{p+1} || z_l - J_\ast v_l ||_F v_l^T \right). 
\]

(4.7)

Similarly to the proof of Lemma 4.1, we obtain the estimate

\[
\left( \sum_{l=1}^{p+1} || z_l - J_\ast v_l ||_F v_l^T \right) \leq C(|| e_{i+1} || + || e_i ||),
\]

(4.8)

where \( e_i \) denotes \( || x_i - x_\ast || \). Hence, using (4.7) and (4.8),

\[
|| E_{i+1} ||_F \leq || E_i \left( I - \frac{s_i s_i^T}{s_i^T s_i} \right) ||_F + C(|| e_{i+1} || + || e_i ||).
\]

(4.9)

This inequality also holds for Broyden’s method and is used in the proof for the \( q \)-superlinear convergence of the method. Using (4.9) in the appropriate place in the proof of Theorem 8.2.2 in [4], the proof also applies to the BSI2 and BSI3 variants of the BSI rank \( p + 1 \) method.
Table 5.1
The dimensionless parameter values for the reverse flow reactor.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_1$</td>
<td>$6.9393 \cdot 10^{-4}$</td>
</tr>
<tr>
<td>$K_2$</td>
<td>0.1749</td>
</tr>
<tr>
<td>$K_3$</td>
<td>$1.5577 \cdot 10^{-6}$</td>
</tr>
<tr>
<td>$K_4$</td>
<td>variable</td>
</tr>
<tr>
<td>$K_5$</td>
<td>$2.4038 \cdot 10^{-3}$</td>
</tr>
<tr>
<td>$K_6$</td>
<td>174.06</td>
</tr>
<tr>
<td>$K_7$</td>
<td>0.01</td>
</tr>
<tr>
<td>$g(\theta)$</td>
<td>$1.6656 \cdot 10^{-5} e^{25.785(\theta-1)/\theta}$</td>
</tr>
<tr>
<td></td>
<td>$1.6656 \cdot 10^{-5} + e^{-25.785/\theta}$</td>
</tr>
</tbody>
</table>

5. The cooled reverse flow reactor. A reverse flow reactor is a packed bed reactor in which the flow direction is periodically reversed to trap a hot zone within the reactor. In this way exothermic reactions can be operated without preheating the feed stream. We consider a cooled reverse flow reactor in which a single irreversible, exothermic, first order reaction occurs. We describe the reverse flow reactor by a one-dimensional, pseudohomogeneous model that accounts for axial heat and mass dispersion and for external mass transfer resistance between the fluid and the catalyst. The basic model assumption is that all the physical properties are independent of the temperature and the concentration. The model we will discuss next is taken from [8].

The variables in the model are the dimensionless temperature $\theta(t,x):[0,\infty) \times [0,1] \to [0,\infty)$ and the conversion $\chi(t,x):[0,\infty) \times [0,1] \to [0,1]$. The dimensionless parameters that appear in the model equations are denoted by $K_j$ for $j = 1, \ldots, 7$ and by $g(\theta(t,x))$. The model-specific values are given in Table 5.1. The parameter $K_4$ will be used as the bifurcation parameter. For the flow from left to right the dimensionless energy and species balances read

\begin{align}
\theta_t &= K_1 \theta_{xx} - K_2 \theta_x + K_3 g(\theta)(1 - \chi) + K_4 (1 - \theta), \\
\chi_t &= K_5 \chi_{xx} - K_6 \chi_x + K_7 g(\theta)(1 - \chi),
\end{align}

with boundary conditions

\begin{align}
K_1 \theta_x(t,0) &= K_2 (\theta(t,0) - 1), \\
K_5 \chi_x(t,0) &= K_6 \chi(t,0), \\
\theta_x(t,1) &= 0, \\
\chi_x(t,1) &= 0.
\end{align}

At each integer value of $t$ the flow direction reverses and the evolution equations and the boundary conditions change accordingly. A cycle consists of two flow reversals. This means that one cycle has a duration of two time units. Equations (5.1)–(5.2) with the boundary conditions (5.3) describe the model.

We would like to compute periodic solutions of (5.1)–(5.2) with the boundary conditions (5.3) that satisfy

\begin{align}
\theta(0,x) &= \theta(1,1-x) \quad \text{and} \quad \chi(0,x) = \chi(1,1-x).
\end{align}

Thus we are interested in periodic solutions that consist of two symmetric parts of one time unit. There might of course also exist periodic solutions without this symmetry, and also periodic solutions with period lengths equal to a multiple of the forcing period. Here we will not compute these solutions, but in principle these solutions can also be computed with the methods discussed in this paper by choosing another period map $F$ (integrating over more than one forcing period).
6. Results and comparison of the methods. For the reverse flow reactor we chose to discretize the spatial variable on a grid of 60 nodes using a finite volume approach with upwinding of the first space derivatives in the mass balance. The discretization of the model equations results, for the reverse flow reactor, in a system of 120 ordinary differential equations. The components in the finite difference discretization can be ordered in such a way that the Jacobian of the system of ordinary differential equations has a band structure. This system of ordinary differential equations is integrated in time using the NAG FORTRAN library routine D02EJF. In this section we present the results for the different methods used to compute a branch of periodic states of the reverse flow reactor. The branch is depicted in Figure 6.1. Here stable periodic states are represented by solid lines, and unstable states are represented by dashed lines. The different bifurcation points are denoted in the figure and explained in the caption.

We used three different values of $p$, i.e., 5, 7, and 9, for all the methods. For the CNP and the CNPGS methods, we also varied the parameter $l$. All the methods used the same initial solution on the branch, and the same initial basis $V_0$. The initial solution and initial basis were computed beforehand.

For all methods we used the same simple step size control procedure. After a convergence failure, the step size is decreased and divided by two, and when a point is computed sufficiently fast, the step size is increased and multiplied by 1.6 (up to a maximum step size). This variable step size strategy allows us to observe not only differences in convergence speed, but also differences in the size of the domain of attraction of the different methods. For each point $(x, \lambda)$ computed on the branch, the three largest eigenvalues of $J_F(x, \lambda)$ (according to absolute value) are computed with four digits of accuracy. For this purpose, after the BSI or Newton–Picard iterations, in most cases only one subspace iteration was sufficient.

Note that the projection step in the subspace iteration algorithm is strictly speaking not necessary if we keep the dimension of the subspace spanned by $V$ fixed during the iterations in the BSI rank $p + 1$ method or the Newton–Picard method. This step is intended for the approach in [13, 12]. The projection step aims to find the $p$ vectors, out of the $(p + p_e)$-dimensional subspace, that span the subspace determined by the $p$ eigenvectors corresponding to the $p$ largest eigenvalues. The projection step does help.
with the additional refining of the approximations to the three largest eigenvalues after convergence of the BSI rank \( p + 1 \) method, for which subspace iteration on the whole subspace \( V_i \) is used [12].

The matrix-vector products \( J_F(x)v \) that are needed in the subspace iteration and in the Newton–Picard method can be calculated in different ways. A first possibility is to use a finite difference approximation given by

\[
J_F(x)v = \frac{\partial F}{\partial x}(x)v \approx \epsilon^{-1}(F(x + \epsilon v, \lambda) - F(x, \lambda)),
\]

where \( (x, \lambda) = x \). This approach costs one extra evaluation of \( F \). Another possibility is to solve the initial value problem

\[
\dot{z}(t) = \frac{\partial f}{\partial x}(t, x(t, x_0), \lambda)z(t), \text{ with } z(0) = v,
\]

where \( (x_0, \lambda) = x \). The matrix-vector product is now given by \( J_F(x)v = z(1) \). This initial value problem is linear, in contrast to the initial value problem that has to be solved when evaluating \( F \). We used both approaches in the tests in this section, for all the methods. We found that if we counted the number of initial value problem solves for each method, there was not much difference in performance between the two approaches. This means that if we write “evaluation of \( F \),” one can also read “initial value problem solve.”

The second, variational approach for computing matrix-vector products has the advantage that with integrators such as ODESSA [11], it is possible to integrate the variational equations concurrently with the time integration for the evaluation of \( F \). With this approach matrix-vector products can be computed cheaply compared to evaluating \( F \). We will focus on this issue when comparing the performance of the BSI rank \( p + 1 \) method with that of Broyden’s method.

The number of evaluations of \( F \) (or number of initial value problem solves) that each method needs to compute the whole branch is listed in Table 6.1. The number of points computed on the curve is listed in Table 6.2.


We first compare the performance of the BSI1 method with the performance of the Newton–Picard methods. We clearly see that the BSI rank \( p + 1 \) method is much more efficient. Actually, the BSI rank \( p + 1 \) method is most efficient for \( p = 5 \), but also for the other values of \( p \) the method is much more efficient than the either the CNP or the CNPGS method.

The next most efficient method is CNP with \( l = 2 \) and \( p = 7 \). Note that there is not much difference between the \( p = 7 \) and \( p = 9 \) cases. For \( p = 5 \), however, the CNP method needs considerably more function evaluations.

The CNPGS method is the least efficient method and needs the most function evaluations. This is clearly due to the fact that it has the smallest domain of attraction. As is seen from Table 6.2, the CNPGS method needs considerably more continuation points than both the BSI and the CNP methods. This fact is also illustrated in Figure 6.2. From this figure, we observe that the BSI rank \( p + 1 \) method uses the maximum step size all along the branch except close to the turning points. The CNP method has more or less the same robustness as the BSI rank \( p + 1 \) method. The CNPGS method is the least robust method, since it is the only method that uses smaller step sizes to compute the unstable part of the branch between the two points where an eigenvalue crosses the unit circle at \(-1\).
Table 6.1

<table>
<thead>
<tr>
<th>Method</th>
<th># $F$ eval. for $p = 5$</th>
<th># $F$ eval. for $p = 7$</th>
<th># $F$ eval. for $p = 9$</th>
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<tr>
<td>BSI1</td>
<td>9242</td>
<td>10000</td>
<td>12255</td>
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<td>9229</td>
<td>10035</td>
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<tr>
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<td>11432</td>
<td>11671</td>
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<td>10374</td>
<td>12774</td>
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<td>11839</td>
<td>12066</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.75$)</td>
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<td>12245</td>
<td>13446</td>
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<tr>
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<td>10704</td>
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<td>BSI2 ($\kappa=0.5$)</td>
<td>11057</td>
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<td>BSI2 ($\kappa=0.75$)</td>
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<td>12639</td>
<td>14423</td>
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<td>BSI3 ($\kappa=0.01$)</td>
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<td>348</td>
<td>370</td>
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<td>BSI3 ($\kappa=0.1$)</td>
<td>361</td>
<td>365</td>
<td>349</td>
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<tr>
<td>BSI3 ($\kappa=0.25$)</td>
<td>360</td>
<td>348</td>
<td>365</td>
</tr>
<tr>
<td>BSI3 ($\kappa=0.5$)</td>
<td>348</td>
<td>365</td>
<td>349</td>
</tr>
<tr>
<td>BSI3 ($\kappa=0.75$)</td>
<td>350</td>
<td>366</td>
<td>362</td>
</tr>
<tr>
<td>CNP ($l=1$)</td>
<td>40296</td>
<td>22268</td>
<td>20900</td>
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<td>CNP ($l=2$)</td>
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<td>19992</td>
<td>20744</td>
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<td>CNP ($l=3$)</td>
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<td>20832</td>
<td>21964</td>
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<td>CNPGS ($l=1$)</td>
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<td>28428</td>
<td>29032</td>
</tr>
<tr>
<td>CNPGS ($l=2$)</td>
<td>31292</td>
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<td>29623</td>
</tr>
<tr>
<td>CNPGS ($l=3$)</td>
<td>27797</td>
<td>28091</td>
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Table 6.2

<table>
<thead>
<tr>
<th>Method</th>
<th># points for $p = 5$</th>
<th># points for $p = 7$</th>
<th># points for $p = 9$</th>
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<tbody>
<tr>
<td>BSI1</td>
<td>348</td>
<td>348</td>
<td>370</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.01$)</td>
<td>348</td>
<td>348</td>
<td>370</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.1$)</td>
<td>361</td>
<td>365</td>
<td>349</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.25$)</td>
<td>360</td>
<td>348</td>
<td>365</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.5$)</td>
<td>348</td>
<td>365</td>
<td>349</td>
</tr>
<tr>
<td>BSI1 ($\kappa=0.75$)</td>
<td>350</td>
<td>366</td>
<td>362</td>
</tr>
<tr>
<td>BSI2 ($\kappa=0.01$)</td>
<td>348</td>
<td>348</td>
<td>370</td>
</tr>
<tr>
<td>BSI2 ($\kappa=0.1$)</td>
<td>372</td>
<td>348</td>
<td>348</td>
</tr>
<tr>
<td>BSI2 ($\kappa=0.25$)</td>
<td>360</td>
<td>348</td>
<td>360</td>
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<tr>
<td>BSI2 ($\kappa=0.5$)</td>
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<td>350</td>
<td>374</td>
</tr>
<tr>
<td>BSI2 ($\kappa=0.75$)</td>
<td>361</td>
<td>362</td>
<td>351</td>
</tr>
<tr>
<td>BSI3 ($\kappa=0.01$)</td>
<td>348</td>
<td>348</td>
<td>370</td>
</tr>
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<td>BSI3 ($\kappa=0.1$)</td>
<td>372</td>
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<td>348</td>
</tr>
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<td>BSI3 ($\kappa=0.25$)</td>
<td>360</td>
<td>348</td>
<td>360</td>
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<tr>
<td>BSI3 ($\kappa=0.5$)</td>
<td>364</td>
<td>350</td>
<td>374</td>
</tr>
<tr>
<td>BSI3 ($\kappa=0.75$)</td>
<td>361</td>
<td>362</td>
<td>351</td>
</tr>
<tr>
<td>B+SI</td>
<td>352</td>
<td>352</td>
<td>352</td>
</tr>
<tr>
<td>CNP ($l=1$)</td>
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<td>355</td>
<td>359</td>
</tr>
<tr>
<td>CNP ($l=2$)</td>
<td>420</td>
<td>345</td>
<td>359</td>
</tr>
<tr>
<td>CNP ($l=3$)</td>
<td>356</td>
<td>350</td>
<td>360</td>
</tr>
<tr>
<td>CNPGS ($l=1$)</td>
<td>415</td>
<td>411</td>
<td>396</td>
</tr>
<tr>
<td>CNPGS ($l=2$)</td>
<td>419</td>
<td>438</td>
<td>408</td>
</tr>
<tr>
<td>CNPGS ($l=3$)</td>
<td>403</td>
<td>395</td>
<td>409</td>
</tr>
</tbody>
</table>

In [13] it is noted that the performance of the Newton–Picard methods in terms of the number of evaluations of $F$ needed is essentially independent of the number of nodes used in the discretization. In the order to check whether this is also the case for the BSI rank $p + 1$ method, we computed the same branch of solutions using a discretization of model equations on 100 nodes, so that $N = 200$. In this case the BSI rank $p + 1$ method with $p = 7$ needed 10397 evaluations of $F$ to compute the whole branch. This number is close to the number of evaluations of $F$ needed for the discretization on 60 nodes.
The step sizes for the different methods along the branch. Along the x-axis the parameter $K_4$ is plotted, along the y-axis the maximal dimensionless temperature in the bed at flow reversal is plotted, and along the z-axis the step size used to compute the point on the branch is plotted. The maximum step size is equal to one. For each method the most efficient run is used: BSI, $p = 7$; CNP, $l = 2$ and $p = 7$; CNPGS, $l = 3$ and $p = 5$.

Table 6.3
Number of calls to the integration code ODESSA for both Broyden’s method and the BSI rank $p + 1$ method ($p = 7$).

<table>
<thead>
<tr>
<th>Method</th>
<th>Total</th>
<th>$G(\omega) = 0$</th>
<th>Eigenvalues</th>
</tr>
</thead>
<tbody>
<tr>
<td>Broyden</td>
<td>4652</td>
<td>3996</td>
<td>656</td>
</tr>
<tr>
<td>BSI1</td>
<td>1621</td>
<td>1560</td>
<td>61</td>
</tr>
</tbody>
</table>

6.2. Comparison of the BSI method and Broyden’s method. If we compare the number of $F$ evaluations for the BSI method with Broyden’s method followed by subspace iterations for the computation of the largest eigenvalues (see Tables 6.1 and 6.2, lines denoted by B+SI), we see that Broyden’s method is slightly more efficient. Because the numbers for the evaluations of $F$ for Broyden’s method and the BSI rank $p + 1$ method are very similar, it is interesting to compare these two methods more precisely.

The BSI method has one clear advantage over Broyden’s method. This advantage is that in one iteration of the BSI method, we are able to compute all the information needed for the updating of the Jacobian with one call to a time integration code such as ODESSA [11]. In this way the directional derivatives that are needed for the updating of the invariant subspace are computed concurrently with the evaluation of $F$.

The number of calls to ODESSA for both Broyden’s method and the BSI rank $p + 1$ method are given in Table 6.3. We see that the total number of calls for the BSI rank $p + 1$ method is far less than half the number for Broyden’s method. The numbers in the second column denote the number of calls used for the computation of the periodic solutions on the branch. This number is equal to the number of iterations both methods need. We see that the extra rank $p$ update for the BSI rank $p + 1$ method speeds up the convergence of Broyden’s method more than twice. The last column contains the number of calls to ODESSA for the computation of the three largest eigenvalues after the convergence to a periodic solution. We see that the BSI rank $p + 1$ method needs only 61 extra calls.

The fact that the BSI rank $p + 1$ method is able to compute almost all the needed matrix-vector product concurrently with the evaluation of $F$ results in favorable
computation times when compared to Broyden’s method, even though Broyden’s method needs less function evaluations: the timing results for the whole branch are 798 seconds for the BSI method and 899 seconds for Broyden’s method. These timing runs were performed on one node of the Beowulf cluster at SARA Computing and Networking Services, Amsterdam.

6.3. The convergence of the BSI method. In order to analyze the convergence behavior of the BSI rank $p + 1$ method, we plot for the BSI1 method the value of $1/c$ along the computed branch of periodic states of the reverse flow reactor; see Figure 6.3. We see that this quantity does not approach zero, except near the end of the branch, but in this part of the branch the BSI1 method converged in one step up to the desired tolerance. This means that for the largest part of the branch we can choose a value of $\kappa$ such that the BSI2 and BSI3 methods, for which we have proved $q$-superlinear convergence, perform exactly the same steps as the BSI1 method. It is an interesting question whether it is possible to obtain an a priori lower bound for $1/c$ in certain cases. To answer this it might be necessary to use the extra structure in the problem: the subspace $V$ is always contained in the $N$-dimensional subspace spanned by the first $N$ standard basis vectors of $\mathbb{R}^{N+1}$ in which the BSI method operates, while the update $s_i$ can be any vector in $\mathbb{R}^{N+1}$. Using this fact and the dependence of the map $F$ on the parameter $\lambda$, it might be possible to obtain an a priori lower bound for $1/c$.

In Tables 6.1 and 6.2 we give the results for the variants BSI2 and BSI3. We see that there is not much difference between the results for the BSI1 variant. Notice that the methods perform better if $\kappa$ is smaller, i.e., if the method is “closer” to the BSI1 method.

7. Conclusions. In this paper we have introduced a Broyden rank $p + 1$ update continuation method with subspace iteration (the BSI rank $p + 1$ method) for the computation of branches of periodic solutions of periodically forced partial differential equations. The method makes efficient use of the ideas behind Broyden’s method and behind the Newton–Picard methods developed by [13]. As the name suggests, the method combines a Broyden method with a subspace iteration procedure for the computation of invariant subspaces. Simultaneously, approximations to a periodic solution and to the eigenvalues that determine the stability are computed. In this way bifurcation points can be detected accurately and efficiently.
We have used the method to compute a branch of periodic states of a cooled reverse flow reactor (see Figure 6.1). We have also used two variants of the Newton–Picard method [13] to compute the same branch. For the dynamical system describing the cooled reverse flow reactor, the BSI rank $p + 1$ method proved to be the most efficient method in terms of number of period map evaluations. In fact, the BSI method needed approximately half the number of evaluations of $F$ as the next most efficient method (CNP with $l = 2$ and $p = 7$). The main advantage of the BSI rank $p + 1$ method, when compared to the Newton–Picard methods, is the small number of period map evaluations per iteration of the method. The method proved also to have the largest domain of attraction. This results in the largest average step size, so that the method needed the fewest continuation points.

Of the two variants of the Newton–Picard method, the CNP and the CNPGS variants, the CNPGS is the least expensive in terms of the number of evaluations of $F$ per iteration. The CNPGS method, however, is also the least robust method of the two, and this results in the poorest overall performance.

We did not use a state-of-the-art implementation of the variants of the Newton–Picard method. Such an implementation—which incorporates locking and deflation in the subspace iteration, adaptive control of the dimension of the subspace, and adaptive control of the number of Picard iterations [12]—probably will perform better in tests than the implementation used in this paper. We believe that the BSI rank $p + 1$ method will also benefit from a more sophisticated variant of the subspace iteration algorithm. How large the gain in efficiency will be is a topic for future investigation.

We also compared the BSI rank $p + 1$ method to Broyden’s method followed by subspace iterations for the computation of the largest eigenvalues. The BSI rank $p + 1$ method and Broyden’s method needed a comparable number of function evaluations for the test problem. When for the matrix-vector products the variational equations are solved (such as is done in the ODESSA code), the BSI rank $p + 1$ exploits the possibilities of such a code better than Broyden’s method. This resulted in shorter CPU times for the BSI rank $p + 1$ method when compared to CPU times for Broyden’s method.

We have proven that the variants BSI2 and BSI3 of the BSI rank $p + 1$ method enjoy q-superlinear convergence. We also showed that, as long as the angle between the update and the approximate invariant subspace stays bounded away from zero, there exist values of $\kappa$ such that the BSI2 and BSI3 methods behave exactly as the BSI1 method. For the presented sample problem, we could experimentally show that this angle between the update and the approximate invariant subspace stays away from zero for the BSI1 method in the crucial parts of the computations.

We experimentally showed that the performance of the BSI rank $p + 1$ method in terms of number of evaluations of $F$ is essentially independent of the number of nodes used in the discretization. This fact is already known for the Newton–Picard methods [13].

Acknowledgment. We would like to thank one of the referees for his very detailed report and his useful and constructive remarks.

REFERENCES