HIGH DETAIL STATIONARY OPTIMIZATION MODELS FOR GAS NETWORKS — PART 2: VALIDATION AND RESULTS.

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ABSTRACT. Due to strict regulatory rules in combination with complex nonlinear physics, major gas network operators in Germany and Europe face hard planning problems that call for optimization. In part 1 of this paper we have developed a suitable model hierarchy for that purpose. Here we consider the more practical aspects of modeling. We validate individual model components against a trusted simulation tool, give a structural overview of the model hierarchy, and use its large variety of approximations to devise robust and efficient solution techniques. An extensive computational study demonstrates the suitability of our models and techniques for application in gas network planning.

1. INTRODUCTION

In part one of this paper [18] we have presented detailed models for stationary optimization in gas transport networks, consisting of physical components that are mostly associated with network wide gas dynamics, and of technical components that are usually associated with specific types of network elements. We have also developed smoothing techniques for nonsmooth components in order to apply optimization algorithms that use first and second order derivatives, and we have presented simplifications and approximations of highly complex components to reduce the computational effort when a lower accuracy is acceptable. This leads to a complicated variety of possible combinations of the component models. In this paper we will first provide a structural overview of the possible combinations in Sect. 2. The individual components will then be checked for correctness and accuracy in Sect. 3 by comparing computational results with the results of a commercial gas network simulation software. In Sect. 4 we consider general optimization techniques for computing practically useful and reliable results efficiently. This includes penalty relaxations to obtain additional information when no feasible solutions can be found, and using the variety of simplified and approximated models to construct sequences of warm-started NLPs for tackling highly detailed and nonsmooth problems. Results will be presented in Sect. 5 for a set of hard problem instances on a real-world network, and for a large set of publicly available instances on an artificial but realistic network. An extensive literature review has already been given in part one [18]; a broader review can be found in [9].

In the following subsection we introduce a fundamental application problem that will be referred to throughout the paper.

1.1. Validation of Nominations (NoVa). The validation of nominations is one of the key problems in the day-to-day work of gas network planners. In a nutshell, the NoVa problem is this:

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FIGURE 1. Model aspect graph of resistors (non-isothermal)

Given a transport network and a nomination of all entry and exit load flows, determine whether there is a technically and physically feasible operation of the network that satisfies this nomination.

The details are as follows. Consider a set of booking contracts with entry and exit customers. The booked capacities define maximal load flows that the customers may *nominate* for actual transport. Now, a *nomination* is defined as a complete set of balanced entry and exit load flows together with specific restrictions on gas pressures and with prescribed values for all quality parameters of the supplied gas, like calorific value, etc. The problem of validating such a nomination is to decide whether it is technically and physically feasible, i.e., whether all load flows can be realized. Throughout the paper we only consider the continuous feasibility problem that is obtained after fixing all discrete decisions. For more information on the NoVa problem and on our overall solution approach see, e.g., [9, 16].

2. NLP MODEL VARIANTS

In part one of this paper [18], we have presented component models for the required element types of gas networks and for basic physical phenomena that are relevant to the entire network. The component models often consist of several sub-models describing different physical phenomena or technical processes that we refer to as model *aspects*, and many of these aspects admit a choice among several modeling variants that we refer to as model *concretizations*. This leads to a large variety of possible NLP models, each of which is determined by choosing a complete set of concretizations for all model aspects.

In this section we provide a structural description of the sets of component models, aspects and concretizations, to obtain an overview and better understanding of the complete set of NLP models and their interrelations. This is complicated by the fact that the selection of concretizations for certain *global* aspects determines the sets of choices of other aspects, whereas the remaining selections are mostly independent of each other, even for different network elements of the same type. We will arrange aspects and concretizations in a directed (meta-)graph to formalize these interdependencies, so that every possible NLP model will ultimately correspond to a forest in the meta-graph: a set of trees that satisfy certain properties.



FIGURE 2. Model aspect graph of resistors (isothermal)

To this end, let us first give an overview of the component models and some illustrative examples of model aspects and concretizations. The basic physical phenomena include: gas compressibility, the equation of state for real gas, the heat capacity of gas, the interdependence of changes in gas pressure and temperature (Joule–Thomson effect), and mixing of different gas compositions. The required element types include: nodes (entries, exits, junctions), passive arcs (pipes, resistors, short cuts), and active arcs (compressor groups, control valve stations, valves). Typical model aspects are the pressure drop along a pipe, the fuel consumption of a compressor, or the equation of state. For instance, the aspect "equation of state" has two concretizations, "thermodynamical standard equation" and "Redlich–Kwong". The former depends on another model aspect, "compressibility", which in turn has two concretizations that we use in our NLP models, with details given in part one; further variants can be added as needed.

There are two *alobal* aspects whose concretizations determine sets of choices of other aspects: gas temperature and gas composition, the latter being represented by seven gas quality parameters. In a coarse model we may fix the values of gas temperature and gas quality parameters globally, but if we decide to take into account changes at *some* network elements, then we must compute changes throughout the network. This is the case since both temperature and composition are globally coupled: fixing certain values at some network elements and computing changes at other elements would inevitably lead to model inconsistencies. Taking the changes into account is known as *temperature tracking* and *gas quality tracking*, respectively. Practitioners tend to use it only when necessary as it complicates the models and computations considerably. Note finally that the seven gas quality parameters (molar mass m, calorific value H_c , pseudocritical pressure and temperature p_c, T_c and the coefficients of isobaric molar heat capacity A, B, C) consist of four independent subsets that can be selected for tracking individually, namely $m, H_c, (p_c, T_c)$ and (A, B, C). Tracking has to be enabled if at least one subset is selected, but only the selected subsets appear in the required mixing equations. The two global aspects thus have 2^5 concretizations arising as combinations of "fixed" or "tracking" for the temperature and the four parameter groups.

Next, we start describing the above-mentioned meta-graph to formalize relations between model aspects and concretizations. As before, let the directed graph $G = (\mathbb{V}, \mathbb{A})$ model the gas network with node set \mathbb{V} and arc set \mathbb{A} . We denote model aspect nodes of the meta-graph by $\alpha \in \mathcal{A}$ and concretization nodes by $\gamma \in \mathcal{C}$; their union $\mathcal{A} \cup \mathcal{C}$ defines the node set of the meta-graph. An arc $a = \alpha \gamma$ from α to γ exists iff γ is a concretization of the model aspect α . These arcs in the set $\mathcal{A} \times \mathcal{C}$ are

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FIGURE 3. Model aspect graph of compressors (nonisothermal)

called "can-be-modeled-by"-arcs. For instance, there is an arc from the model aspect node α = "equation of state" to the concretization node γ = "Redlich-Kwong". In addition, the meta-graph also contains arcs from the set $\mathcal{C} \times \mathcal{A}$. These occur for concretizations that depend on other model aspects. For instance, the concretization "thermodynamical standard equation" depends on the sub-aspect "compressibility" (with concretizations "AGA" and "Papay"). We refer to these arcs in the set $\mathcal{C} \times \mathcal{A}$ as "contains-aspect"-arcs. In contrast, there are no arcs in $\mathcal{C} \times \mathcal{C}$ or in $\mathcal{A} \times \mathcal{A}$ since concretizations do not need to be concretized further and there is no need to consider aspects containing sub-aspects. (For instance, considering the specific change in adiabatic enthalpy $H_{\rm ad}$ as an aspect with sub-aspects "compressibility" and "isentropic exponent" would be physically reasonable but would unnecessarily complicate the meta-graph, cf. Figs. 3–4). Thus, every aspect has at least one concretization and every concretization belongs to some aspect, implying that aspect nodes always have nonzero outdegree and concretization nodes always have nonzero indegree. We define *fundamental* model aspects as aspects with indegree zero, and terminal concretizations as concretizations with outdegree zero. Those form the respective sets $\overline{\mathcal{A}} := \{ \alpha \in \mathcal{A} : |\delta_{\alpha}^{-}| = 0 \}$ and $\overline{\mathcal{C}} := \{ \gamma \in \mathcal{C} : |\delta_{\gamma}^{+}| = 0 \}.$



FIGURE 4. Model aspect graph of compressors (isothermal)

In contrast to the two global model aspects, gas temperature and gas composition, we call all other model aspects *local*. The selection of concretizations of local aspects does not influence the sets of choices of other local aspects (except for subaspects of the concretizations, of course), and the selections are independent for all individual network elements. For instance, we may choose to concretize the compressibility factor model with the AGA formula at pipes in regional (low-pressure) sub-networks but choose to model the compressibility factor with the Papay formula on large (high-pressure) transport pipelines. Similar considerations may apply to the choice between "approximation" or "discretization" of the governing ODEs for pipes. We thus speak of *global* and *local* selections of concretizations.

To select a specific NLP model variant, we first have to fix the global selections and subsequently the local selections for all nodes and arcs of the network graph. Let us denote the global selections by $\sigma_g \in \Sigma_g$ where Σ_g is the (finite) set of global choices. Then, for every network element $\ell \in \mathbb{V} \cup \mathbb{A}$, the global selections σ_g determine a *local model aspect graph* $\mathcal{M}_{\ell}(\sigma_g)$ describing all possible aspects, concretizations and sub-aspects of ℓ ; see, e.g., Figs. 1–4 for the local model aspect graphs of resistors and compressors. Fixing the local selections for ℓ means that, for every fundamental aspect node $\bar{\alpha}$ in $\mathcal{M}_{\ell}(\sigma_g)$, we select a tree $T_{\ell,\bar{\alpha}}$ rooted in $\bar{\alpha}$ such that every aspect node α in $T_{\ell,\bar{\alpha}}$ has outdegree one (select one concretization of α) whereas every concretization node γ in $T_{\ell,\bar{\alpha}}$ has the same outdegree as in $\mathcal{M}_{\ell}(\sigma_g)$ (cover every sub-aspect of γ). Obtaining trees here requires that identical aspects reached from several concretizations are separate nodes in $\mathcal{M}_{\ell}(\sigma_g)$, such as

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"compressibility" in Figs. 1, 3, and 4. Thus $\mathcal{M}_{\ell}(\sigma_g)$ is actually a forest, and every tree $T_{\ell,\bar{\alpha}}$ is a subtree of the tree rooted in $\bar{\alpha}$. On the other hand, we always select identical concretizations for every instance of a "multiple" aspect, and we do not see any good reason to do otherwise. (Therefore we depict every such aspect as a single node with a visual indication of its multiplicity.) Now, the union $\bigcup_{\bar{\alpha}} T_{\ell,\bar{\alpha}}$ is a forest in $\mathcal{M}_{\ell}(\sigma_g)$ representing a specific local model selection. We denote those local selections by $\sigma_{\ell} \in \Sigma_{\ell}(\sigma_g)$ where $\Sigma_{\ell}(\sigma_g)$ is the set of choices (local forests) at element ℓ determined by σ_g . Given network elements ℓ_1, ℓ_2 of the same type, the local model aspect graphs $\mathcal{M}_{\ell_1}(\sigma_g), \mathcal{M}_{\ell_2}(\sigma_g)$ are clearly identical whereas the local selections $\sigma_{\ell_1}, \sigma_{\ell_2}$ may differ. A formal definition of the complete set of possible selections (or NLP model variants) thus reads

$$\Sigma := \bigcup_{\sigma_g \in \Sigma_g} \left\lfloor \{\sigma_g\} \times \prod_{\ell \in \mathbb{V} \cup \mathbb{A}} \Sigma_\ell(\sigma_g) \right\rfloor.$$

The elements of Σ will be denoted by $\sigma = (\sigma_g, \sigma_{\mathbb{V} \cup \mathbb{A}})$ with $\sigma_{\mathbb{V} \cup \mathbb{A}} = (\sigma_\ell)_{\ell \in \mathbb{V} \cup \mathbb{A}}$; each corresponds to a forest in the meta-graph constructed as a union of local forests. The respective subsets of global concretizations and of local concretizations at element $\ell \in \mathbb{V} \cup \mathbb{A}$ selected by σ_g and σ_ℓ will be denoted by $\mathcal{C}_g, \mathcal{C}_\ell$.

With a selection $\sigma \in \Sigma$ at hand, we can now formulate the corresponding NLP model variant of the complete network. To this end, let $\bar{\mathcal{E}}_g, \bar{\mathcal{I}}_g$ and $\bar{\mathcal{E}}_\ell, \bar{\mathcal{I}}_\ell$ denote complete index sets of equality and inequality constraints that may appear in concretizations of the global aspects and of the aspects of element $\ell \in \mathbb{V} \cup \mathbb{A}$, respectively. Likewise, let $\bar{\mathcal{V}}_g, \bar{\mathcal{V}}_\ell$ denote the respective index sets of variables that appear in these concretizations. Here we require that corresponding constraint index sets of different elements $\ell_1, \ell_2 \in \mathbb{V} \cup \mathbb{A}$ are mutually disjoint, even if ℓ_1, ℓ_2 are elements of the same type,

$$\bar{\mathcal{E}}_{\ell_1} \cap \bar{\mathcal{E}}_{\ell_2} = \emptyset, \qquad \bar{\mathcal{I}}_{\ell_1} \cap \bar{\mathcal{I}}_{\ell_2} = \emptyset.$$

In addition, we require that every local variable index is associated with a unique network element ℓ . This is possible since variables of arc $\ell \in \mathbb{A}$ can only appear in the constraints of arc ℓ and of its head and tail nodes whereas variables of node $\ell \in \mathbb{V}$ can only appear in the constraints of node ℓ and of its incident arcs. Thus we obtain the following complete sets of constraints and variables that may appear in the NLP model:

$$\bar{\mathcal{E}} = \bar{\mathcal{E}}_g \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \bar{\mathcal{E}}_\ell, \qquad \bar{\mathcal{I}} = \bar{\mathcal{I}}_g \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \bar{\mathcal{I}}_\ell, \qquad \bar{\mathcal{V}} = \bar{\mathcal{V}}_g \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \bar{\mathcal{V}}_\ell$$

Now, every model selection σ determines certain global and local index subsets corresponding to the constraints and variables of the concretizations C_q, C_ℓ ,

$$\mathcal{E}_{\sigma_g} \subseteq \bar{\mathcal{E}}_g, \quad \mathcal{E}_{\sigma_\ell} \subseteq \bar{\mathcal{E}}_\ell, \qquad \mathcal{I}_{\sigma_g} \subseteq \bar{\mathcal{I}}_g, \quad \mathcal{I}_{\sigma_\ell} \subseteq \bar{\mathcal{I}}_\ell, \qquad \mathcal{V}_{\sigma_g} \subseteq \bar{\mathcal{V}}_g, \quad \mathcal{V}_{\sigma_\ell} \subseteq \bar{\mathcal{V}}_\ell.$$

The unions of these sets yield the complete index subsets associated with $\sigma \in \Sigma$,

$$\mathcal{E}_{\sigma} = \mathcal{E}_{\sigma_g} \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \mathcal{E}_{\sigma_\ell}, \qquad \mathcal{I}_{\sigma} = \mathcal{I}_{\sigma_g} \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \mathcal{I}_{\sigma_\ell}, \qquad \mathcal{V}_{\sigma} = \mathcal{V}_{\sigma_g} \cup \bigcup_{\ell \in \mathbb{V} \cup \mathbb{A}} \mathcal{V}_{\sigma_\ell}.$$

The selected NLP variant finally reads

$$\min_{x_{\sigma}} \quad f(x_{\sigma}) \quad \text{s.t.} \quad c_{\mathcal{E}_{\sigma}}(x_{\sigma}) = 0, \quad c_{\mathcal{I}_{\sigma}}(x_{\sigma}) \ge 0, \tag{1}$$

where

$$x_{\sigma} := (x_i)_{i \in \mathcal{V}_{\sigma}}, \quad c_{\mathcal{E}_{\sigma}} := (c_i)_{i \in \mathcal{E}_{\sigma}}, \quad c_{\mathcal{I}_{\sigma}} := (c_i)_{i \in \mathcal{I}_{\sigma}}.$$

The objective function f, although not part of the model variant, is restricted by its set of variables: it can be any smooth function that depends only on x_{σ} .

We remark that some of the NLP model variants contain additional continuous choices that we will not formalize here. Those appear in concretizations that involve smoothing techniques or discretizations of differential equations, with smoothing parameters or grid parameters that vary in continuous sets. Examples of such concretizations include "Darcy–Weisbach-type smoothed" for resistors (see Fig. 1), or the concretization "discretization" of the model aspect "Momentum equation" for pipes (see Fig. 29 in App. A).

Figs. 1–4 and Fig. 22–29 in App. A show the local model aspect graphs for resistors, compressors, pipes, control valves and nodes. As already mentioned, the given sets of concretizations correspond to the model variants developed in part one of this paper [18]; further variants are conceivable and may be added as needed.

Let us finally remark that the occurrence of model aspects with several concretizations is not restricted to the case of gas network optimization. Our experience shows that other problems from engineering and physics share similar properties. The framework formalized in this section can thus be used in other fields of applied optimization as well.

3. Model Validation

For a proper validation of the model components developed in the first part of this paper [18], one would ideally set up inverse problems to perform parameter estimations based on measurements from a real gas network. Although possible in principle, this is prohibitively expensive in practice. Instead, we validate our model components against a commercial simulation software by a systematic comparison. This is a natural approach since our industrial partners rely on the same simulation software to check the results of the optimization methods.

For a stationary simulation, one generally has to prescribe a suitable set of quantities so that the resulting system of nonlinear differential and algebraic equations has a unique solution. In our case this includes discrete and continuous control settings of active network elements, flows at all but one entry and exit nodes in every connected component of the network, and in addition the pressure at one node in every hydraulically connected area. The simulation problem is typically solved by a Newton framework using numerical integrators. In practical planning applications, the solution is then manually checked against inequality constraints like pressure limits at nodes or the velocity limit on pipes.

Modern simulation methods can compute solutions with high accuracy. For increased computational speed, some simulation tools offer model simplifications, such as replacing heat dynamics by a constant temperature [23] or deactivating the tracking of gas quality parameters like molar mass or calorific value.

To analyze the physical and technical accuracy of our NLP model components, we now carry out a consistency test against the commercial gas network simulation software SIMONE v5.73 [12, 19]. The relevant components are pipes, resistors, control valves, and compressors: entries, exits, junctions, valves, and short cuts have trivial models and are therefore excluded. For the comparison, each of the four element types is tested with several technical and physical settings, and possibly with several model variants. The variant offering the highest common level of detail between the NLP model and SIMONE is always included in the test set. In specific, we use a non-isothermal model of gas physics with gas mixing at nodes, discretized ODEs for the gas flow in pipes, and maximally detailed models of compressors and drives; see [18, Sect. 3] for the details of the NLP model.

When generating the test sets, we aim at choosing realistic parameters of network elements. Ideally, we would like to use the elements of a real-world network for the comparison, but due to limitations of the SIMONE API [13], some parameters can only be entered manually via the graphical user interface, which is impractical for our large number of tests. The test set of every network element type is therefore

Quantity	Tested values	Unit
L	0.01, 0.1, 0.9, 46.0	km
D	150, 310, 405, 1185	mm
k	0.006, 0.02, 0.1, 0.5	mm
h_{out}	-500, 0, 500	m
$p_{ m in}$	3.9, 15.3, 53.8, 74.4	bar
$T_{ m in}$	288.15, 298.15, 308.15, 318.15	Κ
Q_0	50, 250, 500, 750	$1000{ m Nm^3/h}$

TABLE 1. Parameters for test cases of pipes

designed as follows. For every technical parameter (except pipe inclination, which cannot be set with the SIMONE API) we take four quantiles from the distribution of values of a real-world gas network: the 10%, 35%, 65%, and 90% quantiles. The gas network considered is the northern high-calorific network of our industry partner Open Grid Europe¹ (see Sect. 5 for more details of this network). All other parameters of our test sets like pressure or flow values are suitably chosen from typical ranges based on our experience. The Cartesian product of the resulting sets of parameters then defines the test set. Excluding the lower and upper 10% of parameter values helps to avoid unrealistic parameter combinations that would otherwise distort the analysis. We also exclude those cases from the analysis where SIMONE or the NLP solver do not converge or where SIMONE yields a solution that violates the velocity limit on a pipe. The remaining successful cases will be referred to as *valid* tests.

As measures of deviation for a physical or technical quantity x (such as pressure or temperature) we will consider the absolute and relative deviations of x itself,

$$d_{\rm abs}(x) = |x_{\rm NLP} - x_{\rm Sim}|, \qquad d_{\rm rel}(x) = \frac{|x_{\rm NLP} - x_{\rm Sim}|}{|x_{\rm Sim}|},$$

and, if applicable, the relative deviation of the change of x along a network arc,

$$d_{\rm rel}(\Delta x), \quad \Delta x = x_{\rm in} - x_{\rm out}.$$

Here, x_{Sim} and x_{NLP} denote the respective solution values of SIMONE and the NLP model. To obtain a meaningful comparison, the component parameters are always identical in the NLP and in SIMONE, similarly the transport situations (typically defined by Q_0 , p_{in} , T_{in}), and the model variants are selected to agree as well as possible. To avoid that exceptional cases dominate the comparison, we will discuss average deviations rather than maximum deviations.

3.1. **Pipes.** For the validation of pipes we select four model variants: the combinations of two pressure loss models (quadratic approximation and ODE discretization) with two models of the compressibility factor (AGA formula and Papay's formula). For each of these four variants, 12 288 test cases are obtained as combinations of the parameters given in Table 1. The geodesic height of the inflow node is fixed at 0 m, so that 8192 physically possible test cases remain (with $|h_{out}| < L$). The valid tests are used to calculate average deviations of $x \in \{p_{out}, T_{out}\}$. Here $d_{rel}(\Delta x)$ is only computed if the change Δx is at least 1% of the inflow value, to avoid numerical noise dominating in cases where the change is insignificant and hence the denominator Δx becomes very small, which could bias the comparison substantially. Table 2 shows the resulting deviations. As expected, the deviations between the discretized ODE model and SIMONE are smaller than in case of the quadratic approximation.

¹https://www.open-grid-europe.com



TABLE 2. Mean deviations of outflow pressure (p_{out}, bar) and outflow temperature (T_{out}, K) of pipes

FIGURE 5. Logarithmic histograms of outflow pressure and temperature for the model variants "ODE discretization" and "Papay's formula"

Note that SIMONE always applies an implicit integration over time for the Euler equations of gas dynamics [10, 23]. In the stationary case, the PDE solution approaches the solution of the spatial ODE asymptotically, and SIMONE simply uses a large time interval to obtain a highly accurate solution. Nevertheless, the absolute deviation of pressure with the quadratic approximation is only about 0.1 bar, which appears to be sufficiently small for practical purposes in mid- to long-term planning tasks. Figure 5 shows logarithmically scaled histograms of the absolute deviations of outflow pressure and temperature for the model using the discretized ODE and Papay's formula. For the outflow pressure the leftmost bin contains 2574 of 2659 samples (97%) and for outflow temperature the leftmost bin contains 2330 of 2659 samples (88%).

Let us now discuss a specific test instance in more detail. Figure 6 shows some exemplary pressure profiles (outflow pressure vs. flow) of SIMONE and two NLP model variants, each combined with Papay's formula. As one would expect, the outflow pressure deviations between the models grow with increasing pressure loss. In fact, almost arbitrarily large deviations can be generated if the pipe parameters and boundary values are chosen such that huge pressure losses result: for instance, a gap of about 38.5 bar between SIMONE and the quadratic approximation is obtained for the example in Table 3. In this (totally unrealistic) case, the pipe is rather long and its inner wall is extremely rough.

Turning to the outflow temperatures in Figure 7, we observe that all three profiles are roughly comparable but visibly different. An interesting aspect is the difference between the NLP models based on approximation and ODE discretization. As can



FIGURE 6. Outflow pressure profile of an exemplary pipe ($L = 46 \text{ km}, D = 1185 \text{ mm}, k = 0.006 \text{ mm}, s = 0.01, p_{\text{in}} = 74.44 \text{ bar}, T_{\text{in}} = 318.15 \text{ K}$ and $T_{\text{soil}} = 284.15 \text{ K}$) computed using Papay's formula in SIMONE v5.73 (—), NLP with ODE discretization (---), and NLP with a quadratic approximation (…). Figure on the right is a zoom of the dashed frame in the left figure.



FIGURE 7. Outflow temperature profile for the pipe of Fig. 6

TABLE 3. Extremal pipe example

Parameters	Results
$\begin{array}{l} L = 46 \ {\rm km} \\ D = 850 \ {\rm mm} \\ k = 1 \ {\rm mm} \\ s = 0.01 \\ Q_0 = 1944 \times 10^3 \ {\rm Nm^3/h} \\ p_{\rm in} = 199.1 \ {\rm bar} \end{array}$	$p_{\text{out}}(\text{SIMONE}) = 4 \text{ bar}$ $p_{\text{out}}(\text{NLP, ODE discr.}) = 30.6 \text{ bar}$ $p_{\text{out}}(\text{NLP, quadr. approx.}) = 42.5 \text{ bar}$

be seen in Figure 7, the approximation model does not catch the convex curvature at small flows. Nevertheless, the average absolute deviation of 1 K (cf. Table 2) lies within the range of data accuracy: environmental forecasts, soil temperatures, and heat transfer coefficients of pipe walls are inherently inaccurate, and data imprecision is likely to introduce larger errors than 1 K.

Figure 7 also shows a strange behavior of SIMONE for flow values close to zero: there is a sharp peak in the outflow temperature profile that cannot actually represent physical behavior. Therefore we tested the example of Figure 7 again with the more recent software release SIMONE v5.83. This version appears to fix the problem at small flows, see Figure 8 (right). However, both versions exhibit a discontinuity of the outflow pressure at zero flow, see Figure 8 (left). Moreover, comparing both

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FIGURE 8. Differences between SIMONE v5.73 (---) and SI-MONE v5.83 (----) versions for small flows

TABLE 4. Parameters for test cases of control valves

Quantity	Tested values	Unit
Δp	0, 5, 10	bar
$p_{ m in}$	20, 40, 60	bar
$T_{\rm in}$	283.15, 300.15, 318.15	Κ
Q_0	500, 1000	$1000{ m Nm^3/h}$

TABLE 5. Parameters for test cases of resistors. Resistors with a linear pressure loss model are described by the pressure drop ξ and those with a nonlinear (Darcy–Weisbach type) pressure loss model are described by the drag factor ζ and the diameter D.

Quantity	Tested values	Unit
$\begin{array}{c} p_{\mathrm{in}} \\ T_{\mathrm{in}} \\ Q_0 \end{array}$	20, 40, 60 283.15, 300.15, 318.15 500, 1000	$egin{arrsymbol{bar}\ K\ 1000Nm^3/h\ \end{array}$
ξ	2, 4, 6, 8, 10	bar
$\zeta \\ D$	5, 20, 50, 70, 90 300, 525, 775, 1000	1 mm

versions at large flows in the example of Figure 8 yields deviations up to 1.5 bar for outflow pressure and up to 0.93 K for outflow temperature. These deviations are larger than most of the deviations between the NLP model and SIMONE v5.73. Thus, regarding temperature, the NLP model appears to match SIMONE as well as the two versions of SIMONE match each other.

Finally, although the quadratic pressure loss approximation is a substantial simplification of the PDE model solved by SIMONE, its results appear to be sufficiently accurate for practical use in mid- to long-term planning tasks: the deviations discussed above are mostly smaller than data inaccuracies caused by uncertainty of measurements and unknown network data. In any case, if higher physical accuracy is required, the approximation model can easily be replaced by the ODE model.

3.2. Control Valves and Resistors. In order to obtain unique solutions for control valves and resistors, the values of inflow pressure, inflow temperature and flow

Model variant	Valid tests	Quantity	$d_{\rm abs}(x)$	$d_{\rm rel}(x)$	$d_{\rm rel}(\Delta x)$
AGA Papay	$54\\54$	$T_{ m out} \ T_{ m out}$	$0.7390 \\ 0.0938$	$0.0025 \\ 0.0003$	$0.3380 \\ 0.0409$

TABLE 6. Outflow temperature deviations of control valves

TABLE 7. Outflow pressure and temperature deviations of resistors: linear pressure loss model (above) and Darcy–Weisbach type model (below)

Model variant	Valid tests	Quantity	$d_{\rm abs}(x)$	$d_{\rm rel}(x)$	$d_{\rm rel}(\Delta x)$
AGA Papay	90 90	$T_{ m out} \ T_{ m out}$	$0.8850 \\ 0.0983$	$0.0030 \\ 0.0003$	$0.3370 \\ 0.0357$
AGA Papay AGA Papay	317 318 317 318	$p_{ m out}$ $p_{ m out}$ $T_{ m out}$	$\begin{array}{c} 0.0060 \\ 0.0083 \\ 0.6830 \\ 0.2280 \end{array}$	$\begin{array}{c} 0.0006 \\ 0.0008 \\ 0.0024 \\ 0.0008 \end{array}$	$\begin{array}{c} 0.0013 \\ 0.0018 \\ 0.3410 \\ 0.0740 \end{array}$

TABLE 8. Parameters for test cases of piston and turbo compressors

Quantity	Tested values	Unit
Δp	10, 15, 20	bar
$p_{ m in}$	45, 50, 55	bar
$T_{ m in}$	283.15, 300.15, 318.15	Κ
Q_0 (piston compressors)	300, 400, 500	$1000{ m Nm^3/h}$
Q_0 (turbo compressors)	400, 500, 600	$1000{ m Nm^3/h}$

are fixed, and in case of control valves also the pressure reduction. For the compressibility factor we consider again both the AGA formula and Papay's formula. The temperature change in control valves and resistors is calculated by the same model, [18, Eq. (100)], hence similar behavior is to be expected in the comparisons. The definition of the test set is given in Tables 4 and 5 and the results can be seen in Tables 6 and 7. In one respect they are similar to the case of pipes: Papay's formula yields very small deviations for the outflow temperature whereas the AGA formula yields deviations up to 8 times as large. However, the absolute deviations are always reasonably small at less than 1 K.

For control values and for resistors with a linear pressure loss model, the outflow pressure depends linearly on the preset values by [18, Eq. (99)] and [18, Eq. (12)]. Thus the NLP model and SIMONE should agree almost within machine precision, but actually the outflow pressures show an average absolute deviation of about 5×10^{-5} bar. Since the NLP results can be shown to be correct, the deviation possibly indicates an output precision of the SIMONE API of about 1×10^{-4} . The precision of internal computations is not known for SIMONE.

For resistors with a Darcy–Weisbach type pressure loss model, the outflow pressure is determined by the nonlinear equation [18, Eq. (13)]. Although all values in this equation are given (except for the outflow pressure), we observe again small but measurable deviations between the NLP and SIMONE. The mean absolute deviation is 0.006 bar with the AGA formula and 0.008 bar with Papay's formula.

Compressibility factor	Valid tests	Quantity	$d_{\rm abs}(x)$	$d_{\rm rel}(x)$	$d_{\rm rel}(\Delta x)$
AGA	444	$H_{\rm ad}$	0.0729	0.0019	
Papay	435	$H_{\rm ad}$	0.0388	0.0010	
AGA	444	P	0.0058	0.0009	
Papay	435	P	0.0105	0.0016	
AGA	444	T_{out}	0.6810	0.0021	0.0199
Papay	435	$T_{ m out}$	0.6200	0.0019	0.0188
AGA	513	$H_{\rm ad}$	0.0721	0.0021	
Papay	513	$H_{\rm ad}$	0.0437	0.0012	
AGA	513	P	0.0044	0.0014	
Papay	513	P	0.0053	0.0017	
AGA	513	T_{out}	0.6600	0.0021	0.0263
Papay	513	$T_{ m out}$	0.6060	0.0019	0.0243

TABLE 9. Results of the comparison of turbo compressors (above) and piston compressors (below)

3.3. Compressors. For the examination of turbo and piston compressors, we fix the values of inflow pressure, inflow temperature, outflow pressure and flow (see Table 8 for the specific values). The test set is constructed from 18 combinations of physical models of the compressibility factor, the isentropic exponent and the temperature rise equations, yielding 1458 test cases both for turbo and piston compressors. We compare computed values of the specific change in adiabatic enthalpy $H_{\rm ad}$ (kJ kg⁻¹), required compressor power P (MW) and compressor outflow temperature $T_{\rm out}$ (K).

Table 9 shows the results for turbo and piston compressors. Since $H_{\rm ad}$ and P do not have inflow and outflow values, $d_{\rm rel}(\Delta x)$ does not exist for these quantities. The results of the NLP model and SIMONE agree very well for the compressors: the relative deviations $d_{\rm rel}(x)$ are no larger than 0.2%, and even the relative deviations $d_{\rm rel}(\Delta x)$ with respect to the temperature change are just about 2%. A possible reason for the remaining small temperature difference may lie in the model of the isentropic exponent or in differences of the adiabatic efficiency. These two quantities cannot be accessed with the SIMONE API and are therefore not compared.

3.4. **Conclusion.** Of course, one cannot expect that computational results of any simulation software and of the NLP model agree with high precision. Differences in numerical algorithms, discretizations and implementations make deviations almost inevitable. In addition, it is likely that there are differences in data handling and in the precision of data input and output. Obviously, these aspects may add to the deviations caused by the numerical schemes.

As we have seen, it is difficult to validate our highly accurate NLP model against a closed-source simulation package like SIMONE because the code of the latter cannot be inspected. This is the reason why in certain cases we can only observe but not analyze differences in the computational results. For instance, we have no clue why the outflow temperatures of several network element types show unexpectedly large deviations if we use the AGA formula for the compressibility factor, whereas deviations are very small with Papay's formula. A possible reason might be that the correction term for the heat capacity [18, Eq. (38)] is handled in different ways. As stated in part 1 of this paper, that term evaluates to zero with the AGA formula, so the value is *exact* in our NLP model. However, it is possible that SIMONE evaluates the integral numerically rather than simply dropping it. Differences like those may cause many of the unexplained deviations, but we have no way to tell.

In summary, the observed differences between the NLP model and SIMONE are surprisingly small. Thus, we view the comparison presented above as a successful validation of our model components for the intended use in optimization-based mid-term to long-term planning in gas networks.

4. Optimization Techniques

This section addresses two types of techniques that are important when solving the problem of validation of nominations in practice: penalty-based relaxations in several standard and problem-specific variants and sequences of simplified NLPs that are used to obtain a good initial estimate for the NLP of interest.

As it is the case for the framework of model aspect graphs in Sect. 2, the techniques presented in this section are not restricted to the field of gas network optimization but can also be applied to other problems from engineering or physics.

4.1. **Penalty Formulations.** Penalty formulations provide an essential tool for the modeler as well as for the practitioner who uses optimization software to solve real-world problems. For the modeler it is often difficult to find model errors or inconsistencies, particularly if he or she is not also a practitioner or an expert in the area of application. More importantly, the modeler (or practitioner) is often troubled with erroneous, incomplete or inconsistent data. These data problems may lead to infeasible instantiations of the optimization model and are typically hard to detect. Sophisticated penalty formulations relax certain constraints of the optimization model, thus giving a chance to detect the area of the transport network in which a model or data problem may be located. For the practitioner, penalty formulations are also very useful in situations where he or she is not only interested in *optimal* solutions but also in other *feasible* solutions.

Furthermore, penalty formulations are often instrumental in speeding up the solution process by finding an (almost) feasible solution first. It is usually easier and faster to solve the original problem with such a near-feasible solution as initial estimate than solving it from scratch (cf. Sect. 4.2).

In what follows, we present some penalty-based relaxation schemes that are used in gas transport applications. The presented relaxation schemes will be used in Sect. 4.2, and numerical results are finally given in Sect. 5.

4.1.1. Relaxation Schemes for Gas Network Optimization Models. Suppose that a specific instantiation of interest of the model hierarchy presented in [18, Sect. 3] is given in standard NLP form,

$$\min f(x) \quad \text{s.t.} \quad c_{\mathcal{E}}(x) = 0, \quad c_{\mathcal{I}}(x) \ge 0. \tag{2}$$

Here and in what follows, \mathcal{E} and \mathcal{I} are the respective index sets of equality and inequality constraints. With slack variables $s = (s_{\mathcal{E}}^+, s_{\mathcal{E}}^-, s_{\mathcal{I}}^+)$, the associated standard ℓ_1 penalty model is then defined as

$$\min_{x,s} \|s\|_1 \quad \text{s.t.} \quad c_{\mathcal{E}}(x) + s_{\mathcal{E}}^+ - s_{\mathcal{E}}^- = 0, \quad c_{\mathcal{I}}(x) + s_{\mathcal{I}}^+ \ge 0, \quad s \ge 0.$$
(3)

Likewise, the standard ℓ_{∞} penalty model reads

$$\min_{x,s} \|s\|_{\infty} \quad \text{s.t.} \quad c_{\mathcal{E}}(x) + s_{\mathcal{E}}^+ - s_{\mathcal{E}}^- = 0, \quad c_{\mathcal{I}}(x) + s_{\mathcal{I}}^+ \ge 0, \quad s \ge 0.$$
(4)

Both models are full relaxations of (2) and hence feasible by construction. The ℓ_1 penalty model (3) is actually smooth since $||s||_1$ is just the sum of the components

of s, while the ℓ_{∞} penalty model is easily converted to an equivalent smooth formulation by adding an extra variable z for $||s||_{\infty}$ and inequality constraints $z - s_i \ge 0$ for all components of s.

Minimizing the ℓ_1 norm has the well-known property that it tends to generate *sparse* solutions, i.e., solutions with only a few nonzero entries in *s* (see [4, 22] and the references therein). This property is very useful in detecting and analyzing potential reasons of infeasibility.

Minimizing the ℓ_{∞} norm is useful if one is interested in determining the smallest possible upper bound on the constraint violations. In contrast to (3), solutions of (4) tend to have the property that many components of s do not vanish but often have significantly smaller values than the nonzero components of solutions to the corresponding ℓ_1 penalty model.

These two basic relaxation types are now used to develop problem-specific relaxation schemes. In practice, one often does not wish to relax all constraints but only certain classes. For instance, such a class may contain all pressure loss modeling equations on pipes or all constraints modeling the borders of characteristic diagrams of compressors. For handling selected classes of constraints, we introduce the index set $\mathcal{R} \subseteq \mathcal{E} \cup \mathcal{I}$ indicating which constraints are to be relaxed. The required slack variables are $s = (s_{\mathcal{E}\cap\mathcal{R}}^+, s_{\mathcal{E}\cap\mathcal{R}}^-, s_{\mathcal{I}\cap\mathcal{R}}^+)$. With this notation, the partly relaxed ℓ_1 penalty model is given by

$$\min_{x,s} \|s\|_{1} \quad \text{s.t.} \quad c_{\mathcal{E}\setminus\mathcal{R}}(x) = 0, \quad c_{\mathcal{E}\cap\mathcal{R}}(x) + s^{+}_{\mathcal{E}\cap\mathcal{R}} - s^{-}_{\mathcal{E}\cap\mathcal{R}} = 0, \\ c_{\mathcal{I}\setminus\mathcal{R}}(x) \ge 0, \quad c_{\mathcal{I}\cap\mathcal{R}}(x) + s^{+}_{\mathcal{I}\cap\mathcal{R}} \ge 0, \quad s \ge 0.$$
(5)

The corresponding partly relaxed ℓ_{∞} norm penalty model reads

$$\min_{x,s} \|s\|_{\infty} \quad \text{s.t.} \quad c_{\mathcal{E}\backslash\mathcal{R}}(x) = 0, \quad c_{\mathcal{E}\cap\mathcal{R}}(x) + s^+_{\mathcal{E}\cap\mathcal{R}} - s^-_{\mathcal{E}\cap\mathcal{R}} = 0, \\ c_{\mathcal{I}\backslash\mathcal{R}}(x) \ge 0, \quad c_{\mathcal{I}\cap\mathcal{R}}(x) + s^+_{\mathcal{I}\cap\mathcal{R}} \ge 0, \quad s \ge 0.$$
(6)

Note that these two relaxations are not necessarily feasible, except of course with the choice $\mathcal{R} = \mathcal{E} \cup \mathcal{I}$ where they become identical to the standard penalty models.

The partly relaxed penalty models (5) and (6) are of special interest in practice, because a suitable choice of the index set \mathcal{R} may allow to detect reasons of infeasibility. Of course, from the mathematical viewpoint, there is no such thing as a "reason" of infeasibility: infeasibility is simply caused by a set of incompatible constraints, and in general there may be a variety of possibilities to achieve feasibility by relaxing different subsets of those constraints. In practice, however, different constraints often have different relevance (depending on the specific situation). Then, if the violation of a lower-priority constraint makes the problem otherwise feasible, that lower-priority constraint may be interpreted as "the reason" of infeasibility, and a sufficiently small violation will be tolerated.

Suppose that a practitioner is confronted with an infeasible instance. If, say, a relaxation of compressor group constraints leads to feasibility, the reason might be that a lower pressure bound in a downstream part of the network is too tight. Another reason might be that the operating ranges of certain active compressor units are not sufficiently large for generating an outflow pressure that satisfies the pressure bounds in the downstream network. The opposite situation (too tight upper pressure bounds) may be detected by relaxing control valve constraints that limit the pressure decrease from below. Here, the "reason" depends on the interpretation of the situation by the practitioner, who may decide that the pressure bounds or the borders of the characteristic diagrams of the compressors are more important, respectively. Another relaxation that is often used in practice is to relax the supplied and discharged amounts of flow at entry and exit nodes. Given a solution to this relaxed model with non-vanishing slack variables, the practitioner obtains the

information of how much the contracts with one or more entry or exit customers have to be modified in order to achieve a feasible flow situation.

Going further, it may be useful for practitioners to partition the index set \mathcal{R} into several mutually disjoint constraint classes \mathcal{R}_k , $k \in \mathcal{K}$,

$$\mathcal{R} = \bigcup_{k \in \mathcal{K}} \mathcal{R}_k \subseteq \mathcal{E} \cup \mathcal{I},$$

and allow a specific maximum constraint violation \bar{s}_k for each constraint class under consideration. Letting $s_k = (s_{\mathcal{E}\cap\mathcal{R}_k}^+, s_{\mathcal{E}\cap\mathcal{R}_k}^-, s_{\mathcal{I}\cap\mathcal{R}_k}^+)$ and $s = (s_k)_{k\in\mathcal{K}}$, this leads to the modified ℓ_{∞} feasibility problem

Find
$$(x,s)$$
 s.t. $c_{\mathcal{E}\setminus\mathcal{R}}(x) = 0$, $c_{\mathcal{E}\cap\mathcal{R}}(x) + s^+_{\mathcal{E}\cap\mathcal{R}} - s^-_{\mathcal{E}\cap\mathcal{R}} = 0$,
 $c_{\mathcal{I}\setminus\mathcal{R}}(x) \ge 0$, $c_{\mathcal{I}\cap\mathcal{R}}(x) + s^+_{\mathcal{I}\cap\mathcal{R}} \ge 0$, (7)
 $s \ge 0$, $\bar{s}_k - \|s_k\|_{\infty} \ge 0$ for $k \in \mathcal{K}$.

This refined relaxation plays an important role in the presence of soft constraints (see [9, Chap. 11] for a detailed discussion). For instance, the gas flow can excite vibrations of the pipes, which in turn may generate undesired noise in populated areas and, in serious cases, may even destroy the pipes. As there is no suitable quantitative model for these phenomena, a simple speed limit for the gas flow often serves as a crude practical measure to prevent vibrations. In this situation, the network operator may formulate the goal that the speed bound should be satisfied $(v \leq v^+)$, although he will accept small violations if necessary. So he can define the gas velocity constraints to be one of the sets \mathcal{R}_k and can specify an additional quantity \bar{s}_k that softens the former bound to $v \leq v^+ + \bar{s}_k$. Soft constraints like these often play an important role in real-world problems and can be covered by penalty reformulations like (7).

4.2. Sequential NLP Solving. As one might imagine, industrial users would always like to use the most accurate available model. Unfortunately but naturally, the most detailed and accurate model variants presented in [18, Sect. 3] are the most nonlinear and nonconvex ones. In addition, some of them are nonsmooth, i.e., the standard C^2 assumption is violated unless one incorporates numerically challenging smoothing techniques. All these facts illustrate that it is an ambitious task to solve these models from scratch. Since it is impractical for industrial users of optimization methods to adapt the model or to tune the parameters of the solver for individual problem instances, there is a general need for numerically robust solution techniques.

The model hierarchy presented in [18, Sect. 3] suggests a natural way to achieve this goal. The large variety of NLP model variants allow us to set up *sequences* of NLPs that can be solved successively. The key aspect is to arrange each sequence in a way such that successive NLPs differ only slightly in terms of model size and in increase of nonlinearity and nonconvexity, thus allowing for warm starts.

4.2.1. *NLP Sequences.* We consider finite sequences $(\mathsf{NLP}_k), k \in \mathcal{N}$, that are chosen from the family of NLPs given in (1). Recall that \mathcal{A} is the set of all model aspects where aspect $\alpha \in \mathcal{A}$ has concretizations $\gamma_{\alpha} \in \mathcal{C}_{\alpha}$. With this notation, we require that a sequence of NLPs has increasing *accuracy order*, defined by

 $\mathsf{NLP}_k \leq \mathsf{NLP}_{k+1} : \iff \gamma_{\alpha}^k \leq \gamma_{\alpha}^{k+1} \quad \text{for all } \alpha \in \mathcal{A}, \quad k, k+1 \in \mathcal{N}.$

Here, the notation " $\gamma_{\alpha}^{k} \leq \gamma_{\alpha}^{k+1}$ " means that the concretization γ_{α}^{k} of model aspect α used in NLP_{k} is no more accurate than the concretization γ_{α}^{k+1} used in NLP_{k+1} . The notion of "more accurate" is not always defined in a strict mathematical sense (the Papay formula is considered more accurate than the AGA formula, for instance), but in any case we require that all variables of a model variant are also present in a

"more accurate" variant. Thus, the numbers of NLP variables in a sequence satisfy $n_k \leq n_{k+1}$.

4.2.2. Increasing Robustness and Convergence Acceleration. When solving an NLP sequence we make use of three key ideas that improve the solution process by making it more robust and by accelerating convergence of the individual NLPs:

- (1) In most of the sequences used in practice, at least the first NLP is a penalty formulation of the *target NLP* that we actually want to solve; thus solving the first NLP is analogous to phase 1 of the simplex method. The benefit of this approach is twofold. First, solving a penalty model is much easier than solving the *original* model in many cases, according to our experience. Second, if a given instance is infeasible, the penalty formulation allows for a better analysis of possible reasons (cf. Sect. 4.1). In view of this, similar techniques have also been used in optimization for water networks [1].
- (2) Nonsmooth model aspects appear primarily in the model of gas parameter mixing, because flow directions in the network are initially unknown (cf. [18, Eq. (9)]). When solving an NLP of the sequence that incorporates mixing of gas parameters, we fix the flow directions on all network arcs based on the solution of the previous NLP, as follows. Let $q_a^k \in [q_a^-, q_a^+]$ denote the bounded flow variable of arc a in NLP_k and let $(q_a^k)^*$ denote its optimal value in NLP_k. The flow direction for NLP_{k+1} is then fixed by setting properly restricted bounds,

$$q_a^{k+1} \in \begin{cases} \mathbb{R}_{\geq 0} \cap [q_a^-, q_a^+], & (q_a^k)^* > 0, \\ \mathbb{R}_{\leq 0} \cap [q_a^-, q_a^+], & (q_a^k)^* < 0, \\ \{0\}, & (q_a^k)^* = 0. \end{cases}$$
(8)

Similar ideas can be used for other nonsmooth aspects, like (de-)activating gas coolers [18, Eq. (68)] and preheaters [18, Eq. (103)]. It would be reasonable to consider the optimal values of the dual variables of NLP_k in order to relax the third case in (8), yielding restricted bounds $\mathbb{R}_{\geq 0} \cap [q_a^-, q_a^+]$ or $\mathbb{R}_{\leq 0} \cap [q_a^-, q_a^+]$ instead of fixing the flow to zero. We have not done this in the current study because of the technical effort for accessing the dual variables from many NLP codes with different interfaces that we used, and because some solvers do not even provide dual variables.

(3) We initialize the variables of NLP_{k+1} , $x^{k+1} \in \mathbb{R}^{n_{k+1}}$, based on the optimal solution of NLP_k , $(x^k)^* \in \mathbb{R}^{n_k}$. This is done as follows. After some reordering we can write $x^{k+1} = (x^k, \tilde{x}^{k+1})$ with $\tilde{x}^{k+1} \in \mathbb{R}^{\tilde{n}}$, $\tilde{n} = n_{k+1} - n_k$. The key idea is to fix the x^k part at $(x^k)^*$ and to determine \tilde{x}^{k+1} by simple techniques so that $((x^k)^*, \tilde{x}^{k+1})$ satisfies the constraints of NLP_{k+1} as well as possible. For instance, consider a constraint $c(x^k, \tilde{x}^{k+1}_i) = 0$ that can be solved explicitly for the "new" variable \tilde{x}^{k+1}_i in terms of the "old" variables,

$$0 = c(x^k, \tilde{x}_i^{k+1}) \iff \tilde{x}_i^{k+1} = \tilde{c}(x^k) \eqqcolon \tilde{x}_i^*.$$

In this case we initialize \tilde{x}_i^{k+1} with \tilde{x}_i^* . This technique is successively applied to new variables that depend on x^k and already initialized components \tilde{x}_i^{k+1} . In other situations, we simply initialize the new variables with suitable constants.

4.2.3. Stopping Criterion. In the finite sequences considered thus far, $\mathsf{NLP}_{|\mathcal{N}|}$ is the most accurate model that we attempt to solve. It is also possible to extend the presented technique to infinite sequences of NLPs. For instance, one may wish to increase the physical accuracy of the model of gas dynamics in pipes by discretizing the ODEs on successively finer grids, which calls for a *stopping criterion*. Various

criteria are possible, and a natural requirement is that they should be based on changes in variables that exist in every member NLP_k of the sequence. Meaningful quantities of this type are pressures at nodes and mass flows on arcs of the network. With the vectors

$$p_{\mathbb{V}}^k := (p_i^k)_{i \in \mathbb{V}}, \quad q_{\mathbb{A}}^k := (q_a^k)_{a \in \mathbb{A}}$$

and tolerances $\varepsilon_p, \varepsilon_q > 0$, a suitable stopping criterion is given by

$$\|p_{\mathbb{V}}^{k} - p_{\mathbb{V}}^{k-1}\|_{\infty} < \varepsilon_{p} \quad \text{and} \quad \|q_{\mathbb{A}}^{k} - q_{\mathbb{A}}^{k-1}\|_{\infty} < \varepsilon_{q}.$$

$$\tag{9}$$

4.2.4. Specific NLP Sequences. In this section we give three examples of sequences that are useful in gas network optimization. The first sequence mainly addresses the nonsmoothness of the gas quality parameter mixing model given in [18, Sect. 3.3]. The second one deals with highly detailed modeling of gas dynamics in pipes and the last one can be used in order to incorporate heat dynamics in the problem. Computational results of the sequential NLP approach are presented in Sect. 5. For all sequences, we assume that all discrete decisions are given.

An NLP Sequence for Gas Parameter Tracking. The finite sequence for gas quality parameter tracking is informally defined as follows:

- NLP₁: Simple mass conservation model at nodes without gas parameter mixing [18, Eq. (7)], isothermal quadratic approximation of gas dynamics in pipes [18, Eq. (55)], full isothermal compressor group model with machine models as described in [18, Sect. 3.4.10] and [18, Sect. 3.4.15], isothermal control valve model [18, Eq. (99)], standard resistor, short cut and valve models, arbitrary penalty formulation (see Sect. 4.1). Globally constant gas density under normal conditions.
- NLP₂: Like NLP₁ but with an extended node model incorporating mixing of molar masses [18, Eq. (9)].
- NLP₃: Like NLP₂ but with an extended node model incorporating mixing of calorific values.
- NLP₄: Like NLP₃ but with an extended node model incorporating mixing of pseudocritical pressures and temperatures.
- NLP₅: Like NLP₄ but with gas density under normal conditions depending on the quality parameters.

To obtain smooth formulations of the mixing models it is necessary to fix the flow directions on all network arcs, as already discussed in the previous section.

An NLP Sequence for Highly Detailed Isothermal Gas Dynamics Modeling. In this sequence the main focus is on highly detailed modeling of isothermal gas dynamics in pipes. The sequence is given as follows:

 NLP_1 : Identical to NLP_1 in the sequence above.

NLP₂: Like NLP₁ but with the momentum equation discretized on initial grids Δ_a^0 for all pipes $a \in \mathbb{A}_{pi}$ (cf. [18, Sect. 3.4.7]).

 $\mathsf{NLP}_k, k > 2$: Like NLP_{k-1} but with the momentum equation discretized on refined grids $\Delta_a^{k-2} \supset \Delta_a^{k-3}$ for all pipes $a \in \mathbb{A}_{\mathrm{pi}}$.

The canonical stopping criterion for this infinite sequence is (9). To reduce the numerical effort during iterative grid refinement (k > 2), one might also employ an adaptive discretization scheme. Here *adaptive* means that the discretization on pipe $a = ij \in A_{pi}$ is only refined in NLP_k if

$$\left\| \begin{pmatrix} p_i \\ p_j \end{pmatrix}^k - \begin{pmatrix} p_i \\ p_j \end{pmatrix}^{k-1} \right\|_{\infty} > \varepsilon_p \quad \text{or} \quad |q_a^k - q_a^{k-1}| > \varepsilon_q.$$
(10)

Of course, it is also possible to use pipe-specific tolerances, i.e., to replace ε_p and ε_q in (10) by $\varepsilon_{p,a}$ and $\varepsilon_{q,a}$.

An NLP Sequence for Temperature Dynamics Modeling. Here we incorporate all temperature dynamics constraints in the model. The sequence is defined as follows: NLP_1 : Identical to NLP_1 in the sequences above.

- NLP_2 : Like NLP_1 but with an extended node model incorporating the mixing of the coefficients A, B, C of the molar heat capacity of ideal gas.
- NLP₃: Like NLP₂ but with non-isothermal models of all elements of the network (including heat capacity models).

Notice that none of the three sequences above have a cost minimization objective: all of them solve penalty-based relaxation models. However, all sequences can readily be extended or modified to incorporate a cost minimization objective.

5. Computational Results

In order to show the practical relevance of our NLP models and solution techniques on large-scale real-world networks, we now present an extensive computational study for the problem of validation of nominations (NoVa). A complete NoVa model contains discrete aspects of controllable elements as well as detailed models of gas dynamics and technical devices of the network. This combination leads to hard mixed-integer nonlinear optimization (or feasibility) models that are intractable for state-of-the-art general-purpose MINLP solvers on real-world network sizes. In the research project ForNe we have developed custom solution approaches for NoVa. The main idea is to split up the solution process into two stages. The first stage solves a model containing all discrete aspects and simplified variants of the underlying gas physics and technical network elements. Afterwards, the resulting discrete controls are fixed throughout the network in order to obtain a purely continuous and highly detailed NLP model that "validates" the fixed decisions. In this stage, penalty formulations as described in Sect. 4.1 are used to prove feasibility (or ε feasibility) of the underlying MINLP, or to obtain information on possible reasons of infeasibility otherwise. Here we use the sequential NLP (SNLP) approach for solving NLP models with a high level of physical and technical accuracy.

This section deals with that second stage of the NoVa solution approach and presents detailed numerical results for it. We will indeed see that the NoVa problem is solvable even with high detail and accuracy, and that the SNLP approach is a key tool for increasing robustness. Additional information concerning the outcomes of the ForNe project and extensive computational studies of the overall approach are given in [5, 14, 16] and in the book [9].

The remainder of this section is organized as follows. In Sect. 5.1 we present the computational setup including the description of used modeling languages, solvers, and other software and hardware issues. Section 5.2 then describes both the real-world network and the publicly available academic network that we used in our computational experiments. The following Sects. 5.3 and 5.4 present and discuss the respective numerical results on the real-world and the academic test set. Finally, Sect. 5.5 concludes with a discussion of the results.

5.1. Computational Setup. All models and algorithms are implemented using the framework LaMaTTO++ for modeling and solving mixed-integer nonlinear optimization problems on networks [11]. The computational study of the NLP validation stage presented in this paper is run on a desktop PC with a six-core AMD Opteron Processor 2435 with 2600 MHz and 64 G RAM. The operating system is Debian 7.5 and the C++ code LaMaTTO++ is compiled using GCC 4.7.2. All models are implemented in GAMS 24.1.3 [6, 17].

The computational results presented below are obtained using the NLP solver CONOPT 3.15L, because it performs best on the given problem class according to

	HN	GasLib-582
Element type	Number	Number
vertex	747	582
entry	30	31
exit	139	129
junction	578	422
arc	786	609
pipe	505	278
compressor group	8	5
control valve	27	23
valve	50	26
resistor	37	8
short cut	159	269

TABLE 10. Elements of the HN network and the GasLib-582 network

TABLE 11. Exit states of direct approach and of SNLP sequence (HN test set, "parameter tracking")

Exit state	Direct	SNLP
locally optimal	10	30
locally infeasible	19	0
intermediate nonoptimal	1	0

our experience gained during the ForNe project. The solvers we have tested include lpopt [21], CONOPT and CONOPT4 [3], KNITRO [2], SNOPT [8] and MINOS [15].

5.2. Test Instances. We evaluate and test our models on two different test sets based on country-sized transport networks. The first test set, called HN, contains 30 difficult expert instances that arise at OGE. The corresponding network is the northern high-calorific gas network of OGE. The second test set, called GasLib-582, is publicly available [7] and is approximately of the same size as the HN network. These two test sets appear under the same names in other publications as well. Table 10 gives the numbers of elements of the two networks. In order to limit the computational effort to a reasonable amount, we randomly chose 500 out of the 4227 GasLib-582 nominations that are freely available at http://gaslib.zib.de. The first stage solvers of our NoVa approach obtained feasible solutions for 394 of the 500 random nominations. We perform the NLP validation on these 394 instances; see Table 15 in App. C for the complete list.

The results on the HN test set will demonstrate the practicability of our NLP models on hard instances that arise in the day-to-day work at german gas network planning departments whereas the GasLib-582 test set provides publicly available instances on which other researchers can test and compare their algorithms.

5.3. The HN Test Set. Here we present the computational results for the SNLP sequences "parameter tracking", "temperature dynamics" and "ODE discretization" (see Sect. 4.2) applied to the expert instances on the HN test set, and compare them with the *direct approach*, i.e., the attempt to solve the final NLP of each sequence (the *target NLP*) from scratch. SNLP results are determined by the target NLP (exit state, accuracy) or by the entire sequence (iteration count, computing time).



FIGURE 9. Constraint violations of direct approach and of SNLP sequence (HN test set, "parameter tracking").



FIGURE 10. Iteration counts and computing times (s) of direct approach and of SNLP sequence (HN test set, "parameter tracking")

5.3.1. *Parameter Tracking Problem.* Recall that we expect that solving a sequence of NLPs while successively adding new gas parameters to the model leads to better convergence than the direct approach. This is indeed the case, as can be seen in Table 11: the exit states show that the direct approach solves only one third of all instances to local optimality whereas the SNLP approach solves every instance.

Since the NoVa problem is a feasibility problem, our objective is to minimize the constraint violation (using the relaxed NLP in Sect. 4.1). Figure 9 presents empirical distribution functions of the respective objective values for the SNLP sequence and the direct approach, giving for every constraint violation the percentage of instances not exceeding that value. The plot clearly shows that the direct approach is much less successful: it yields substantially larger constraint violations and in fact many "false infeasibles". Moreover, only for 2 out of the 30 instances, it computes a zero constraint violation where the SNLP approach fails to do so.

Figure 10 shows the empirical distribution functions for total iteration counts and computing times (i.e., accumulated over all NLPs in case of the SNLP sequence). Despite the fact that the SNLP approach solves five NLPs rather than just one, both approaches require roughly the same effort. While the SNLP approach needs a slightly larger number of iterations than the direct approach, it does not possess as extreme outliers as the latter. In summary, it can be clearly stated that the SNLP



FIGURE 11. Iteration counts (left) and computing times (s, right) of NLPs of the SNLP sequence (HN test set, "parameter tracking")

TABLE 12. Exit states of direct approach and of SNLP sequence (HN test set, "temperature dynamics")

Exit state	Direct	SNLP
locally optimal	8	30
locally infeasible	6	0
intermediate nonoptimal	1	0
intermediate infeasible	15	0

approach is much more effective than the direct approach in producing feasible solutions while the computational cost is comparable.

The above observations are confirmed by Fig. 11, which displays the distributions of iteration counts and computing times of individual NLPs in the sequence over the HN test set. Except for a few outliers, all iteration counts are below 100, which is approximately 1/5 of the average iteration count of the direct approach. Outliers above 100 are only present in the first, third and fifth NLP of the sequence. Possible explanations are as follows. The first model does not contain any mixing aspects, and the outliers thus may arise from "correcting" the continuous part of the solution given by the first stage of the NoVa solution approach. The third NLP adds the calorific value as a new gas quality parameter to the model. Since the HN test set includes heat power bounds at the entries and exits (in addition to the mass flow bounds), activating the tracking of the calorific value plays a crucial role for the feasibility of the problem. Finally, the fifth and last NLP adds the gas density under normal conditions as a variable to the model. Since this quantity occurs in many constraints—especially in all flow conversion constraints—the worst-case difficulty of the model increases, which might be the reason for outliers in SNLP iteration 5. Turning to the computing times, we see that the majority are below 4s, and we observe a similar qualitative behavior as for the iteration counts.

Let us now consider some model statistics of the NLPs of the SNLP sequence. In all NLPs the number of variables is only slightly larger than the number of constraints: after fixing the discrete decisions only a few degrees of freedom remain, resulting from continuous control variables of active network elements. Both the numbers of variables and constraints strictly increase within the SNLP sequence, lying between 5000 and 15 000. The number of non-constant entries and the total number of entries in the constraints Jacobian also strictly increase, lying between



FIGURE 12. Constraint violations of direct approach and of SNLP sequence (HN test set, "temperature dynamics")



FIGURE 13. Iteration counts and computing times (s) of direct approach and of SNLP sequence (HN test set, "temp. dynamics")

 $10\,000$ and $20\,000$ and between $20\,000$ and $40\,000,$ respectively. The fraction of non-zero entries in the Jacobian is approximately $0.025\,\%.$

5.3.2. Temperature Dynamics Problem. The results for "temperature dynamics" are similar to the results for "parameter tracking". Again, the direct approach solves only an unsatisfactory share of instances to local optimality (26.67%) whereas the SNLP approach solves all instances (see Table 12). Moreover, the SNLP approach reduces the constraint violation much stronger than the direct approach, as Fig. 12 clearly shows. Only two instances are solved with a smaller constraint violation by the direct approach than by the SNLP approach. Furthermore, the SNLP approach is substantially more efficient. Figure 13 compares iteration counts and computing times using empirical distribution functions. On average, the SNLP approach is faster and does not possess as strong outliers as the direct approach does. Iteration counts of the individual NLPs of the SNLP sequence are displayed in Fig. 14. In particular, the first NLP (isothermal model) appears to be the problem with the hardest instances of the sequence, as in "parameter tracking": the number of outliers of NLP 1 clearly exceeds the numbers of outliers of NLPs 2 and 3.

5.3.3. *ODE Discretization Problem*. As a third example of a useful SNLP sequence, we finally discuss the numerical results for "ODE discretization". For this sequence, we always compute five NLPs, which corresponds to three steps of grid refinement



FIGURE 14. Iteration counts (left) and computing times (s, right) of NLPs of the SNLP sequence (HN test set, "temp. dynamics")

TABLE 13. Exit states of direct approach and of SNLP approach (HN test set, "ODE discretization")



FIGURE 15. Constraint violations of direct approach and of SNLP approach (HN test set, "ODE discretization")

(cf. Sect. 4.2.4). Here the results strongly differ from the results of the "parameter tracking" and "temperature dynamics" problems. As Table 13 shows, the direct approach is much more successful on this problem type than for the two problems discussed so far: it solves 90% of all 30 HN instances to local optimality. The SNLP approach again solves all of the instances. In addition, the direct approach has only four outliers with respect to constraint violations (cf. Fig. 15), while the SNLP approach only fails at one instance in finding a locally optimal solution with negligible constraint violation. All other instances are solved by both approaches to local optimality with a vanishing objective (i.e., with zero constraint violation). Thus, speaking only in terms of robustness, the SNLP approach does not offer any major advantage for these instances. However, it turns out that it again outperforms the direct approach significantly with respect to iteration counts and computing

25



FIGURE 16. Iteration counts and computing times (s) of direct approach and of SNLP sequence (HN test set, "ODE discr.")



FIGURE 17. Iteration counts of NLPs of the SNLP sequence (HN test set, "ODE discretization")

times (see Fig. 16). The iteration counts of the direct approach vary between a small number and more than 8000 iterations (corresponding to a computing time of more than 200 s) whereas the SNLP approach almost always requires less than 1000 iterations in total (corresponding to less than 25 s computing time).

Iteration counts for every NLP of the SNLP sequence are given in Fig. 17.

5.4. The GasLib-582 Test Set. Next we describe the results for the three SNLP sequences "parameter tracking", "temperature dynamics" and "ODE discretization" on the random subset of all GasLib-582 instances given in Table 15 of App. C.

5.4.1. Parameter Tracking and Temperature Dynamics Problems. The characteristics of the computational results for "parameter tracking" and "temperature dynamics" differ significantly from the corresponding results on the HN test set. While it is very hard to compute feasible solutions with the direct approach on the latter (cf. Sect. 5.3), this is not the case for the GasLib-582 test set. Table 14 shows the associated exit states for all sequences. The percentage of instances solved to local optimality by the direct approach is 96.4% for "parameter tracking" and 95.7% for "temperature dynamics". By way of comparison, the SNLP approach solves 98.2% and 98.5% to local optimality, respectively. This shows that the SNLP approach is again more robust, but this time only by a small margin. In contrast to the HN test set there are now also a few instances (7 for "parameter tracking" and 6 for

	"paran	n. tracking"	"temp.	dynamics"	"ODE	discr."
Exit state	Direct	SNLP	Direct	SNLP	Direct	SNLP
locally optimal	380	387	377	388	360	371
broken sequence	_	7	_	6	_	22
locally infeasible	14	0	15	0	33	1
intermediate nonopt.	0	0	2	0	1	0
10	0%					
10						
7	5%		 +			
5	0%		i i 1			
5						

TABLE 14. Exit states of direct approach and of SNLP sequence (GasLib-582 test set, all three problem types)

FIGURE 18. Constraint violations of direct approach and of SNLP sequence (GasLib-582 test set, "parameter tracking")

150

Constraint violation

100

SNLP approach

250

300

-- Direct approach

200

25%

0%

0

50

"temperature dynamics") on which the SNLP approach fails completely because one NLP of the corresponding sequence cannot be solved at all (see the row "broken sequence" in Table 14). All other results of these two problems look very similar to the corresponding results on the HN test set. We only discuss the results for "parameter tracking" in detail—all figures for "temperature dynamics" are given in App. B. As Fig. 18 and 19 show, both approaches behave quite similar in terms of constraint violations, iteration counts and computing times. The advantage of the SNLP approach is that it does not possess extreme outliers. However, these outliers of the direct approach are not as extreme as for the HN test set. The qualitative behavior of individual iteration counts and numbers of variables, constraints, etc., of the SNLP approach is similar to the behavior on the HN test set and therefore not further discussed here.

5.4.2. ODE Discretization Problem. For "ODE discretization", both approaches behave similar with respect to the number of instances that are solved to local optimality. The direct approach solves 91.4% of the test set, the SNLP approach is slightly more robust with 94.2%. This is similar to the corresponding results on the HN test set. However, now the SNLP approach has 5.5% broken sequences, which is significantly higher than for the other problems. Except for some outliers in the direct approach, both approaches are also comparable with respect to constraint violations (see Fig. 20). Nevertheless, the SNLP is again clearly preferable since it outperforms the direct approach significantly in terms of iteration counts and computing times (see Fig. 21).

5.5. **Discussion.** In this section we have presented an extensive computational study of our NLP models on two different test sets involving the solution of more



FIGURE 19. Iteration counts and computing times of direct approach and SNLP sequence (GasLib-582 test set, "param. tracking")



FIGURE 20. Constraint violations of direct approach and of SNLP sequence (GasLib-582 test set, "ODE discretization")



FIGURE 21. Iteration counts and computing times of direct approach and of SNLP sequence (GasLib-582 test set, "ODE discr.")

than 6700 NLPs. The NLP models under consideration are hard to solve due to their high degree of nonlinearity and non-convexity and because they possess only a small number of free variables.

It turns out that the expert instances of the HN test set are much harder than the instances of the GasLib-582 test set. Especially the problems in which additional model aspects are successively introduced in the SNLP approach ("parameter tracking" and "temperature dynamics") are hardly solvable from scratch. In these cases, the SNLP approach proves much more robust and entirely practical: it solves all instances of the HN test set successfully, while iteration counts and solution times are comparable to the direct approach. For the "ODE discretization", on the other hand, both approaches are comparable with respect to their robustness. This is to be expected since no additional model aspects are introduced—only the spatial discretization of pipes is refined over the course of the SNLP sequence. However, for the "ODE discretization" problem the SNLP approach clearly outperforms the direct approach on both test sets in terms of iteration counts and solution times.

6. Summary

In part 1 of this paper we describe detailed models for stationary optimization in gas transport networks. This paper provides an analysis of their structural interplay and possible combinations of these models and shows that their level of detail and accuracy is comparable with today's commercial gas network simulation software. Finally, we present tailored optimization techniques and show that—using these techniques—it is possible to solve the resulting NLPs on real-world networks. This demonstrates that optimization models and solvers can provide a valuable tool for gas network planners that adds new capabilities to existing simulation tools.

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Appendix A. Model Aspect Graphs



FIGURE 22. Model aspect graph of nodes (isothermal, mixing of gas types)



FIGURE 23. Model aspect graph of nodes (isothermal, uniform gas composition)



FIGURE 24. Model aspect graph of nodes (non-isothermal, mixing of gas types)



FIGURE 25. Model aspect graph of nodes (non-isothermal, uniform gas composition)



FIGURE 26. Model aspect graph of control valves (isothermal)



FIGURE 27. Model aspect graph of control valves (non-isothermal)

30



FIGURE 28. Model aspect graph of pipes (isothermal). (*) If the exact solutions of the momentum equation are chosen as a concretization, not every choice of the compressibility factor is possible, cf. [18]. In addition, not every combination of pipe slope and compressibility factor is possible. (o) To achieve a smooth NLP model, flow bound strengthening has to be applied. (+) Only possible for horizontal pipes



FIGURE 29. Model aspect graph of pipes (non-isothermal)





FIGURE 30. Constraint violations of direct approach and of SNLP sequence (GasLib-582 test set, "temperature dynamics")



FIGURE 31. Iteration counts and computing times (s) of direct approach and of SNLP sequence (GasLib-582 test set, "temperature dynamics")

Appendix C. GasLib-582 Instances

TABLE 15. ID's of GasLib-582 instances used in the computational study of Sect. 5.4. The set of instances can also be downloaded at www.ifam.uni-hannover.de/~mcs/papers/data.

nomination_freezing_ <id></id>													
35	51	119	121	164	238	362	402	692	714	728	792	831	1023
1093	1106	1135	1191	1233	1263	1416	1505	1568	1588	1599	1775	1889	2005
2055	2110	2145	2206	2236	2293	2351	2378	2466	2491	2567	2629	2724	2969
3044	3078	3126	3225	3383	3465	3496	3500	3592	3673	3853	3910	4028	4182
nomination_cold_ <id></id>													
20	72	276	298	435	486	527	586	595	611	641	681	712	716
796	823	890	893	1033	1056	1132	1156	1309	1311	1327	1363	1380	1432
1439	1461	1466	1468	1543	1689	1703	1748	1765	1792	1828	1851	1910	1951
1971	2038	2067	2077	2124	2147	2182	2189	2192	2195	2311	2316	2406	2425
2438	2478	2486	2510	2516	2551	2556	2627	2686	2763	2791	2799	2857	2891
2898	3064	3086	3136	3155	3202	3387	3395	3397	3407	3528	3539	3543	3566
3620	3624	3707	3769	3800	3824	3878	3987	4038	4105	4129	4146	4176	4200
nomination_cool_ <id></id>													
3	23	89	97	122	141	148	161	201	222	235	277	412	416
456	526	543	565	578	585	628	662	699	724	790	815	821	845
875	879	958	995	1030	1234	1294	1324	1373	1391	1393	1397	1398	1452
1453	1500	1501	1586	1607	1628	1666	1676	1680	1733	1766	1770	1772	1854
1895	1896	1929	1954	2028	2045	2059	2083	2109	2171	2208	2238	2254	2270
2275	2335	2386	2453	2529	2535	2544	2592	2608	2654	2671	2740	2764	2828
2859	2895	2901	2966	3035	3040	3100	3103	3113	3172	3209	3254	3321	3354
3374	3400	3409	3420	3421	3458	3522	3595	3656	3681	3770	3771	3779	3791
3830	3885	3908	3909	3929	3947	3986	4031	4043	4067	4071	4084	4115	4120
4173	4187	4192											
nomination_mild_ <id></id>													
2	18	39	47	92	173	241	246	305	521	630	686	711	749
928	952	1011	1016	1187	1203	1281	1307	1344	1386	1422	1459	1472	1481
1559	1646	1743	1840	1853	1961	1964	2044	2157	2495	2531	2539	2636	2897
2950	3124	3151	3182	3197	3230	3419	3480	3524	3552	3661	3701	3734	3766
3848	3890	4170											
nomination_warm_ <id></id>													
120	136	244	306	441	540	684	786	844	880	916	1062	1080	1215
1521	1523	1556	1656	1747	1803	1812	1975	2060	2140	2221	2259	2276	2314
2356	2399	2584	2646	2648	2689	2718	2824	2988	3048	3050	3164	3235	3331
3341	3501	3718	3742	3749	3784	3956	3962	4160	4168				

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