

Gorash, Y. and Bickley, A. and Gozalo, F. (2018) Assessment of leak tightness for swellable elastomeric seals considering fluid-structure interaction with the CEL approach. In: 24th International Conference on Fluid Sealing, 2018-03-07 - 2018-03-08.

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Assessment of leak tightness for swellable elastomeric seals considering fluid-structure interaction with the CEL approach

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ABSTRACT

Swellable elastomeric seal is a type of specifically engineered packer that swell upon contact with wellbore fluids. Assessment of leakage tightness is a fundamental aspect in the design of swellable packers, since they should guarantee a reliable sealing under extreme pressures of the downhole fluids. Numerical capability of the leakage pressure prediction would facilitate improvement in the packer design methodology. Previous work was focused on investigation of the non-parametric optimisation capability seeking for an optimal external shape with a goal to maximise the grip of a packer with a borehole. The verification of an optimised design was done with a dynamic FE-simulation of packer's failure by extrusion under an excessive pressure. The downside of that verification analysis was that Abaqus/Explicit solver couldn't implement a realistic adaptive pressure application due to changing packer disposition and contact conditions. This simulation challenge is addressed in this paper by application of the Coupled Eulerian-Lagrangian (CEL) approach in Abaqus/Explicit, which provides the ability to simulate a class of problems where the fluid-structure interaction (FSI) is important.

1. INTRODUCTION

Swellable elastomeric seal is a type of specifically engineered packer that swell upon contact with wellbore fluids. Such packers have been widely employed in various oil-&-gas applications including slimming of well design, zonal isolation, water shut-off, and multi-stage fracturing. Assessment of leakage tightness is a fundamental aspect in the design of swellable packers, since they should guarantee a reliable sealing under extreme pressures of the downhole fluids up to 10000 psi (69 MPa). Downhole conditions are difficult to be reproduced using physical testing environment, but feasible to be simulated [1] in virtual environment using FE-codes. Numerical capability of the leakage pressure prediction under different downhole conditions (type of downhole fluid, pressure build-up rate, diameter of the borehole, etc.) would facilitate improvement in the packer design methodology and would allow efficient optimisation of a packer design. Previous work [2, 3] was focused on investigation of the non-parametric optimisation capability seeking for an optimal external shape with a goal to maximise the grip of a packer with a borehole by maximising the contact pressure between them. For this purpose, Tosca/Structure

optimisation suite was used within the Abaqus/CAE environment for maximum computational performance. The verification of an optimised design was done through the dynamic FE-simulation of packer's failure by extrusion under an excessive pressure. The downside of that verification analysis was that Abagus/Explicit solver couldn't implement a realistic adaptive pressure application due to changing packer disposition and contact conditions. This simulation challenge is addressed in this work by application of the Coupled Eulerian-Lagrangian (CEL) approach in Abaqus/Explicit, which provides engineers with the ability to simulate a class of problems where the fluid-structure interaction (FSI) is important, like seals. This capability does not rely on the coupling of multiple software products, but instead solves the FSI simultaneously within single Abaqus environment. The most relevant example of the CEL application to investigation of leakage tightness is the study [4], where the CEL approach predicts not only the pressure at which the seal blows off, but also how the fluid behaves when leakage starts. Apart from the technology demonstration [4], there is a very limited availability of literature sources focused on FSI modelling, that combines extremely large deformations of hyperelastic structures with CEL to address changing contact conditions between fluid and structure.

The idea similar to the one implemented in [4] lies beneath the given numerical study – to investigate a feasibility of FSI simulation with CEL in application to failure analysis of swellable packers. The feasibility assessment would ideally include the computational costs and robustness level of this type of analysis considering the specific conditions including incompressible nature of the material, high pressure applied as a loading and extremely large deformations as a result of excessive pressure application.

In general, the objective of this research project is to develop a design tool integrated into Abaqus/CAE environment to implement the parametric numerical studies using advanced FE-simulation to provide an improved design of packers for various downhole conditions. However, the implementation of the packer's swelling and failure simulations is associated with a number of technical/numerical challenges specific to this particular class of multiphysics problems, which are illustrated in Fig.1 and listed below:

1. Material model. The key component is an advanced material model comprising both hyperelasticity and moisture swelling. It has to consider two-way interaction between mechanical response and swelling capacity. Implementation of such a material model requires using COMSOL Multiphysics [5] or programming of a Fortran subroutine for the user defined material using the Flory & Rehner (1943) theoretical background [6], which is presented in the first instance by Flory-Rehner equation in the following form:

$$-\left[\ln\left(1-\nu_{2}\right)+\nu_{2}+\chi_{1}\nu_{2}^{2}\right]=V_{1}n\left(\nu_{2}^{\frac{1}{3}}-\frac{\nu_{2}}{2}\right),$$
(1)

where v_2 is the volume fraction of polymer in the swollen mass, V_1 is the molar volume of the solvent, *n* is the number of network chain segments bounded on both ends by cross-links, and χ_1 is the Flory solvent-polymer interaction term.

In polymer science Eq.(1) describes the mixing of polymer and liquid molecules as predicted by the equilibrium swelling theory of Flory & Rehner [6]. It describes the equilibrium swelling of a lightly cross-linked polymer in terms of cross-link density and the quality of the solvent. The theory considers forces arising from three sources:

- the entropy change caused by mixing of polymer and solvent;
- the entropy change caused by reduction in number of possible chain conformations via swelling;
- the heat of mixing of polymer and solvent, which may be positive, negative, or zero.

2. Fluid-structure interaction. The moisture swelling process is not uniform and starts on the surfaces which are subject to fluid. Adsorption, which governs the progress of swelling can occur only at free surfaces. Therefore, the fluid pressure penetration needs to be incorporated into the simulation [7] and directly linked to swelling. Distributed pressure penetration load allows for the simulation of fluid penetrating into the surface between two contacting bodies, penetration of fluid from multiple locations on the surface, and application of the fluid pressure normal to the surfaces. It automatically adjusts the application of a fluid pressure depending on changes of contact conditions.

3. Large deformation convergence. Non-uniform swelling is associated with a localised increase of material volume. It may cause a significant distortion of FE mesh and arouse FEA convergence problems. To overcome this, there are a few options available in the setup of the FE-model [8] including a mesh-to-mesh solution mapping (Abaqus/Standard), adaptive remeshing (Abaqus/Explicit) and element distortion control. Convergence issue is crucial to the successful solution of elastomeric structures FE-simulation, because in most cases the FE-analysis fails because of excessive distortion or collapse of elements.

4. Parametric analysis automation. Parametric study assumes considering a large number of different geometric configurations, looking at material properties variation and different downhole conditions. Basically this means a search for an optimal geometry through a sensitivity study, which would result in specific design recommendations for the geometry of a packer. Therefore, it would be reasonable to automate the analysis procedure through an Abaqus plug-in [9] with a convenient graphical user interface (GUI), which provides access to the parameters of geometry, material properties and service conditions.



Figure 1: Diagram of technical requirements for a robust FE-simulation of swellable packers for progressive failure and leakage

2. VALIDATION OF SHAPE OPTIMISATION

In previous works [2, 3] the feasibility of non-parametric optimisation [10] in application to swellable packers was investigated following the successful outcomes of [11] that revealed a great potential of the topology and shape optimization under contact conditions. For that purpose, Simulia Tosca Structure was used – a software system for non-parametric structural optimisation with interfaces to the most of industry standard FE-solvers. Using optimisation techniques, contact pressure in contact zones could be either minimised [12] or maximised as needed in this research. Therefore, the shape optimisation was used [2, 3] to improve the grip of a packer with the surface of a borehole. The normal shape optimization stimulated the surface growth in contact zones, which resulted in a higher contact pressure and shrinkage in a lower. For a trial shape optimisation study, the trimmed version of a packer geometry [1] was used as benchmark problem with *L* reduced from 16" to 2". The optimisation analysis resulted in a rippled external surface of a packer as shown in Fig.2a with comparison to the original rectangular profile. The distribution of contact pressure became very non-uniform as shown in Fig.2b with four maximums, which are about 5 times higher than the original smooth contact pressure.



Figure 2: Shape optimisation of the packer profile with Tosca Structure: a) change of profile geometry and b) corresponding change of contact pressure

An important part of optimisation analysis is a validation of the obtained design, which in this study is expressed in terms of comparative sealing capability. The basic qualitative validation analysis was performed using the general static simulation procedure with implicit solver in Abaqus/Standard [2]. For a more comprehensive and quantitative validation of the packer design, the simulation capabilities of Abaqus/Standard solver were found insufficient. The advantage of Abaqus/Standard implicit solver was a fast solution and the availability of PPL interaction [7]. This functionality replaces the computationally expensive fluid-structure interaction, when the structural analysis has a priority. On other hand, the disadvantage of implicit solver is that the automatic adaptive remeshing is not available as a standard functionality, so the extrusion problems with extreme deformation can't be effectively solved using this product. Therefore, the subsequent work [3] was implemented with the dynamic solver in Abaqus/Explicit, which is recognised as a more robust solver when it comes to very non-linear problems and extremely large deformations.

Abaqus/Explicit was computationally more expensive compared to Abaqus/Standard, but this obstacle was overcome by running simulations on HPC facility. This solver significantly expands the progressive failure analysis capabilities, and actually eliminates any limitations related to non-linearities, large deformations and transient / dynamic effects. The best prove of its efficiency is a solution of a so-called press-fit problem [13], when a cylindrical rubber block compressed from the tube of bigger diameter into the tube with a smaller diameter. In previous work [14] an attempt to develop a robust approach to

simulation has failed. A simple and stable solution for such a benchmark problem using standard implicit solvers in Ansys and Abaqus couldn't be obtained. It should be noted that the successful simulation of press-fit problem [13] became possible only after the modification of a friction model used in analysis from the linear Coulomb to the bi-linear Coulomb-Orowan law [15] expressed in terms of friction force as

$$F_f = \min(\mu | F_n |, F_\tau), \qquad (2)$$

where μ is a coefficient of friction, F_n is a normal force, and F_r is a critical share force, which corresponds to a critical shear stress τ_c in the FEA setup. The Coulomb term $\mu |F_n|$ is linear and describes the partial slip. When the critical value of τ_c is reached, the total slip occurs, which plays a key role in simulation convergence, because it prevents the rubber material from sticking to the relatively rigid walls.

So the work [3] was focused on the development of a practical approach to simulations of packers with Abaqus/Explicit, since the setup of analyses in Standard and Explicit solvers is quite different. The biggest advantages attributed to Explicit solver are automatic adaptive remeshing (in application to large plastic deformations) or distortion control of elements (in application to large hyperelastic deformations) and stable solution of contact problems with large relative displacements. Considering a superior robustness of Abaqus/Explicit, it is a minor drawback that PPL functionality is unavailable for dynamic analysis. The robustness of extrusion failure simulations for swell packers were demonstrated in [3] with advanced validation analysis of the benchmark problem.

Since PPL is unavailable, the pressure was applied to the bottom surface and ramped in the course of simulation for both benchmark packers – original and optimised. The stable and robust convergence has been achieved with the CAX3 element type -a 3-node linear axisymmetric triangle with the activated distortion control having length ratio 0.5. This means that the FE-model topology is adjusted when an element under uniaxial compression undergoes 50% of nominal strain. This FE-mesh adjustment technique [8] together with the bi-linear Coulomb-Orowan friction law provides a guaranteed convergence of a dynamic solution in Abaqus/Explicit. The absence of hourglass issues is provided automatically by the triangular shape of the elements. In should be also noted that in order to accelerate the analysis and facilitate the convergence the default Abaqus/Explicit compressibility ratio (initial bulk modulus to initial shear modulus) $K_0 / \mu_0 = 20$ has been used [3], corresponding to Poisson's ratio v of 0.475. Since typical unfilled elastomers have K_0 / μ_0 ratios in the range of 1,000 to 10,000 (v = 0.4995 to v = 0.49995) and filled elastomers have K_0 / μ_0 ratios in the range of 50 to 200 (v = 0.490 to v = 0.497), this default provides much more compressibility than is available in most elastomers [16]. The forced incompressibility will become more feasible in the future version of ABAQUS (2018) with introduction of the hybrid formulation for elements used in Abaqus/Explicit solver.



Figure 3: FE-meshes of (a) original and (b) optimised benchmark packer geometries



Figure 4: Validation simulation of the benchmark packer failure using original geometry with Abaqus/Standard and mesh distortion control

Comparison of simulation results showed [3] that the optimised packer can bear about 10% of more pressure compared to the original packer with a smooth surface providing and additional validation of optimisation results. The validation simulation of a full-size real packer [1] demonstrated a complete extrusion of the packer [3]. It also showed that extrusion was not gradual, it was rather abrupt with a distinctive critical pressure when sticking to protective rings can't stop progressive slipping, caused by friction and material compressibility. With the recent findings related to convergence facilitation techniques, it was decided to revisit the static implicit simulations with Abaqus/Standard [2] in a view of limited analysis functionality in terms of realistic incompressibility and load application. Figure 3 shows the FE-meshes of (a) original and (b) optimised benchmark packer geometries consisting of the CAX4R element type, 4-node bilinear axisymmetric quadrilateral with reduced integration and distortion control having length ratio 0.5.

The material parameters for van der Waals hyperelastic model were taken from [17] with following values: $\mu = 0.385$, $\lambda_m = 10.35$; $\alpha = 0.279$, $\beta = 0.95$ and D = 0.001. They are based on Treloar's experimental set of stress-strain data for vulcanised rubber [18]. Since not a triangular-shaped element type was used in analysis with incompressible material, it required an additional hourglass control to stabilise its behaviour at very large strains. The stiffness hourglassing control has been used with the stiffness coefficient of 50, which provided a robust convergence for the benchmark packers simulations with results shown

in Figs 4 and 5. In definition of general static step, the automatic adaptive solution stabilisation with specified dissipated energy fraction (0.0002) and maximum ratio of stabilisation to strain energy (0.05) were used. In this case, the combination of Coulomb-Orowan friction law, element distortion control, stiffness hourglassing control and automatic adaptive solution stabilisation helped to achieve a stable simulation of elastomeric component in axisymmetric formulation.



Figure 5: Validation simulation of the benchmark packer failure using optimised geometry with Abaqus/Standard and mesh distortion control

The failure modes of original and optimised packers are significantly different as can be seen from Figs 4c and 5d. The burst pressure in case of optimised packer is also 30% higher than for original one (2.12 MPa vs 3 MPa). The leakage for the original packer occurs in a trivial and predicted way with a fluid pressure burst through the opened contact between packer and borehole as illustrated step-by-step in Fig.4. When pressure builds up, the fluid propagates only in one direction parallel to axis Y. When reaching critical pressure, the packer gets partly extruded through the gap between the protective ring and borehole.

Failure mode of the optimised packer is significantly different and occurs in non-trivial way as illustrated step-by-step in Fig.5. When pressure builds up, the fluid propagates in two directions as specified in Fig.5b - above the packer (through the opened contact between packer and borehole) and underneath the packer (the opened contact between packer and pipe). Contact opening above the packer lost its priority for fluid penetration because of the stronger grip between packer and borehole induced by the rippled packer surface. So it is easier for fluid to propagate in contact opening between packer and pipe. because of less contact pressure and less friction. This scenario results in complete separation of packer from pipe and almost it's complete extrusion through the gap between the protective ring and borehole. The extrusion is progressive and happens quickly almost without increase of pressure, when packer collapses approximately in its middle location and folds. It should be noted that even following the extrusion of the packer, the start of leakage is not explicit. The fluid pressure penetration results in the formation of cavity filled with fluid in the location of packer folding. The cavity just goes on filling with fluid and growing without indication of pressure burst into outer space. The simulated scenario may seem unrealistic, because the packer should fail and rupture before filling with fluid. But this effect can be implemented only with inclusion of progressive material damage.

This numerical simulation finding indicates an interesting structural behaviour effect, which is worth of further investigation, because it may result in a potential design improvement. Regarding the validation analysis of the full-size packer [1], unfortunately it is still not feasible even with recently discovered convergence improvement techniques. Moreover, current and previous studies [2, 3] showed that leakage is not static, it is a rather dynamic process accompanied by the formation of fluid cavities, their expansion and coalescence. Therefore, the dynamic analysis procedure supposed to be more adequate for realistic structural behaviour simulations of full-size packers.

3. FLUID-STRUCTURE INTERACTION

Since in previously conducted dynamic validation simulations with Abaqus/Explicit [3] interaction with fluid was not considered, they are lacking a realism, because the packer failure mode with a leakage through the contact surface can't be modelled. This simulation challenge can be addressed by an application of the Coupled Eulerian-Lagrangian (CEL) approach in Abaqus/Explicit, which provides engineers with the ability to simulate a class of problems where the interaction between structures and fluids is important. This capability does not rely on the coupling of multiple software products, but instead solves the fluid-structure interaction (FSI) simultaneously within the single Abaqus environment [19]. The potential of CEL approach for packers' leakage simulation is investigated below. The highest level of realism in simulation of leakage process is expected from engaging CEL in ABAQUS/Explicit. In order to develop a practical approach to the solution of this class of FSI problems and to understand corresponding capabilities and challenges, a leakage benchmark problem has been developed with the geometry shown in Fig.6 (all

dimension in m). The assembly includes the following components: 1) computational fluid domain; 2) initial fluid volume; 3) rigid stationary walls (top, bottom, back); 4) rigid moving plunger; 5) deformable rubber seal constrained to the bottom wall.







Figure 7: FE-mesh of computational fluid domain and solid parts encapsulated in it

The idea of this benchmark is to build up a fluid pressure by moving the plunger towards the seal in order to induce a progressive deformation of rubber and subsequent leakage through the gap between deformed seal and top wall. All the fluid should be gradually displaced from the left cavity into the right void during the course of simulation. The fluid pressure is monitored during this process, so that the value of a burst pressure is identified by associating it to the moment in time, when the leakage occurs for the first time. It should be noted that for a simplicity the right cavity is considered to be a void in this study. For more realism, a presence of air can be considered in future simulations using separate initial Eulerian volumes and properties definition for a fluid and for a gas. The FE-model shown in Fig.7 comprises the following type of elements:

- 24780 fluid Eulerian FEs (type EC3D8R) / 28080 nodes (water),
- 640 solid Lagrangian FEs (type C3D8R) / 945 nodes (rubber seal),
- 960 rigid shell FEs (type S4R) / 1116 nodes (rigid walls).

Since the number of Eulerian FEs exceeds almost 40 times the number of Lagrangian FEs, it is Eulerian part of the model, that is the most computationally expensive. The fluid element type, EC3D8R – 8-node linear Eulerian brick with reduced integration and hourglass control, is the only available type of fluid FE for CEL. The seal is meshed with C3D8R – 8-node linear brick with reduced integration, distortion control (length ratio 0.1) and enhanced hourglass control, which is quite sufficient to model an incompressible hyperelastic material undergoing moderate deformations. Since the hyperelastic materials parameters are required to be in SI units to avoid compatibility issues with the fluid material model, a new parameters identification has been implemented using internal Abaqus curve fitting tool [20] using Treloar's experimental set [18] with stress in Pa. In terms of strain energy potential, the 3rd order of Ogden form has been used resulting in the fit shown in Fig.8 and the following set of parameters: $\mu_1 = 371784.2$ [Pa], $\alpha_1 = 1.45175$, $\mu_2 = 1308.63$ [Pa], $\alpha_2 = 5.4886$, $\mu_3 = 15445.055$ [Pa], $\alpha_3 = -1.87468$, $D_1 = 5.1477E-10$. The elastic strain observed in the seal is not rally high - just around 6%, when the leakage starts (see Fig.9a), and it goes up to 14% (see Fig.10a), when the fluid flow intensifies, and finally reaches 19% (see Fig.10b), when the relocation of the fluid is finished.

The fluid material properties used in CEL simulation are based on the linear U_s-U_p Hugoniot form of Mie-Grüneisen equation of state model [21], which is used to model compressible viscous and inviscid laminar flow governed by the Navier-Stokes equation of motion. In general, the equation of state is assumed for the pressure as a function of the current density ρ and the internal energy per unit mass E_m as $p = f(\rho, E_m)$, which defines all the equilibrium states that can exist in a material. The internal energy can be eliminated from the above equation to obtain a p versus V relationship (where V is the current volume) or, equivalently, a p versus $1/\rho$ relationship that is unique to the material described by the equation of state model, and it is called the Hugoniot curve. The Hugoniot pressure p_H is a function of density only and can be defined in general from fitting experimental data. An equation of state is linear in energy when it is written in the form:

$$p = f + g E_m, \tag{3}$$

where $f(\rho)$ and $g(\rho)$ are functions of density only and depend on the particular equation of state model. A Mie-Grüneisen equation of state is linear in energy:

$$p - p_H = \Gamma \rho \left(E_m - E_H \right), \tag{4}$$

where p_H and E_H are the Hugoniot pressure and specific energy (per unit mass) and are functions of density only, and $\Gamma = \Gamma_0 \rho_0 / \rho$ is the Grüneisen ratio with Γ_0 as a material constant and ρ_0 as a reference density.

The Hugoniot energy E_H is related to the Hugoniot pressure p_H by

$$E_H = \frac{p_H \eta}{2 \rho_0},\tag{5}$$

where $\eta = 1 - \rho_0 / \rho$ is the nominal volumetric compressive strain. Elimination of Γ and E_H from the above equations (4)-(5) yields

$$p = p_H \left(1 - \frac{\Gamma_0 \eta}{2} \right) + \Gamma_0 \rho_0 E_m .$$
⁽⁶⁾

A common fit to the Hugoniot data is given by

$$p_{H} = \frac{\rho_{0} c_{0}^{2} \eta}{\left(1 - s \eta\right)^{2}},\tag{7}$$

where the speed of sound in a medium c_0 and the dynamic viscosity *s* define the linear relationship between the shock velocity U_s and the particle velocity U_p as follows:

$$U_{s} = c_{0} + s U_{p} \,. \tag{8}$$

With the above assumptions (6)-(8) the linear $U_s - U_p$ Hugoniot form is written as

$$p = \frac{\rho_0 c_0^2 \eta}{\left(1 - s \eta\right)^2} \left(1 - \frac{\Gamma_0 \eta}{2}\right) + \Gamma_0 \rho_0 E_m , \qquad (9)$$

where $\rho_0 c_0^2$ is equivalent to the elastic bulk modulus at small nominal strains. It should be noted that there is a limited amount of compression with a limiting compression given by the denominator of this form of the equation of state $\eta_{\text{lim}} = 1/s$ or $\rho_{\text{lim}} = s \rho_0/(s-1)$.

The definition of a material using the material model (3)-(9) requires c_0 and s, which are in this case taken as a general case corresponding to water at room temperature – $c_0 = 1483$ [m/s] and s = 0.001 [Pa·s] with the density $\rho = 1000$ [kg/m³] for dynamic FEA.



Figure 8: Rubber stress-strains curves fitted with the 3rd order Ogden model

The obtained water pressure distribution is quite noisy spatially as seen from Fig.9b, but it has a random maximum, which can be traced during the simulation. The history of the water pressure maximum (kPa) vs displacement of plunger (mm) is shown in Fig.11, which also displays the real time of simulation (s) on the secondary axis to show the smooth history of displacement that gradually accelerates and stops to reduce the dynamic effects.



Figure 9: The moment of time corresponding to the start of leakage corresponding to (a) maximum principal true strain in seal and (b) pressure distribution in water



Figure 10: True strain in seal during the (a) progression of flow and (b) flow cut-off



Figure 11: Fluid pressure (kPa) vs displacement of plunger (mm) vs time (s)

The observed volumetric (Fig.9b) and history fluctuations (Fig.11) of the liquid can be in the first instance associated with the dynamic type of simulation, then with some compressibility of liquid, and a non-smooth character of the slip of the seal after reaching the critical pressure and associated seal vibrations and waves in the adjacent liquid. Despite of the noisy character of the maximum pressure history in Fig.11, two distinctive global peaks can be recognised - first corresponding to the displacements 9-20 mm and second for 34-40 mm. The second peak is associated with the finishing of water displacement. when pressure goes up because the left cavity gets fully filled and resists to any further water intake. The purpose of this simulation is to obtain the first pressure peak, which indicates the start of leakage. With the displacement of plunger, the water moves in the same direction creating a pressure on the seal. In its turn, seal reacts to this loading by induction of internal stress and corresponding deformation (see Fig.9a). The contact between the seal and top wall is frictional with coefficient of friction of 0.3, that doesn't let the top seal surface to slip easily under growing water pressure. The hyperelastic nature of seal and slipping in contact limits the maximum pressure that is achieved in simulation. A kind of plateau is observed in Fig.11 starting at 9 mm, that indicates the beginning of seal slipping, with the subsequent loss of contact at 15 mm. The noisy history of water pressure is smoothened using the moving average function with 8 periods producing a more clear history of the pressure pulsation illustrated with a solid line in Fig. 11. The averaged signal gives the value of 150 kPa as an indication of the burst pressure. This value is achieved at 9 mm of displacement, that corresponds to 0.33 s of simulation time.

Similar approach is applied to the packers benchmark problems, which are extended from axial symmetry to full 3D considering 1° of rotation as shown in Fig.12 with two elements per thickness of the model and 1 mm of characteristic element dimension. Packer, initial fluid volume, moving plunger and stationary walls are all encapsulated into the global Eulerian computational fluid domain, which has to go beyond the Lagrangian components. The type of FE is exactly the same as was used for CEL benchmark above – C3D8R. But this time it was not sufficient for the converged simulations – both models failed when the fluid and seals got into hard contact and the pressure started to build up as shown in Fig.13. The failures were caused by excessive distortion of the corners elements when contacting with fluid – both distortion and enhanced hourglass control didn't work as it was expected. The elements completely buckled resulting in unrealistic strains and termination of FEA.



Figure 12: Geometries of the benchmark problems: a) original and b) optimised



Figure 13: Partial solutions of the benchmark problems: a) original & b) optimised

4. CONCLUSIONS

The further work will focus on search of a robust FE-model setup, which would guarantee a stable convergence at high fluid pressure in the range from 1 to 100 MPa. It is worthwhile trying the C3D4 type of FE – 4-node linear tetrahedron in combination with distortion control, because in previous work [3] the stable convergence has been achieved with the 3-node linear triangle (CAX3). It should be noted that Abaqus/Explicit simulations with CEL approach are very computationally expensive, e.g. the obtaining the results shown in Fig.13 required around 1000 CPU-hours, and they are still incomplete. Therefore, further work will also focus on improving the computational efficiency of the validation simulations. Static implicit analysis if fast, but not robust, on other hand dynamic explicit analysis is robust, but slow. Therefore, taking the best from both solvers, in the form of dynamic implicit analysis as a form of FSI is available in dynamic solvers and can be applied to large-deformation enclosures (seals, etc.) with adaptive acoustic meshes for fluids [22].

5. ACKNOWLEDGEMENTS

The authors appreciate the financial support of Weir Group PLC within the WARC project "Design Optimisation of Swell Packers" and University of Strathclyde for hosting during the course of this work. Results were obtained using the EPSRC funded ARCHIE-WeSt High Performance Computer (www.archie-west.ac.uk) EPSRC grant no. EP/K000586/1.

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