



OLI Software Release Notes V11

Contents

Introduction.....	3
Version 11.0.1.9.....	4
General Information.....	4
Databank Updates.....	5
Mixed-solvent electrolyte (MSE) databank.....	5
MSE-SRK databank	9
Aqueous corrosion kinetics databank	9
OLI Databook – Bugs & Features.....	13
Engine/Solver - Features.....	13
Engine/Solver - Bugs.....	13
OLI Studio/Analyzer - Features	14
OLI Studio/Analyzer - Bugs	14
OLI Flowsheet: ESP / ESP-FS - Features	17
OLI Flowsheet: ESP / ESP-FS - Bug.....	17
Alliance Products - Features	21
Alliance Products - Bugs	21
OLI Chemistry Wizard (all products).....	21
OLI Developer Edition - General.....	22
OLI Framework - General.....	22

Introduction

For many years, the OLI software release notes were maintained on several different web pages. While this was suitable at the time it has become far too cumbersome to maintain.

With the release of OLI Software V9.6 the release notes have been combined into a single document.

Each version will be listed in reverse chronological order. Within each version update to the products will be listed in this order:

- Databank updates
- Databank fixes
- Engine/solver
- OLI Studio/Analyzer
- OLI Flowsheet: ESP
- OLI Chemistry Wizard
- OLI Developer Edition
- OLI Alliance products
 - Aspen Hysys
 - Aspen Plus
 - gProms
 - IDEAS
 - Petro-Sim
 - Proll
 - Unisim Design
- OLI Security/License Manager

Version 11.0.1.9

General Information

Version 11 introduces the concept of vertical markets. This means we have segmented our databases to reflect these vertical markets. Some species, previously available to all licensees, are now segregated into individual verticals. These “Segments” are now licensed separately.

For existing licenses that exist for the release of version 11, no segments are implemented. Users with existing licenses can use all these species.

The databanks were released on the same date as the first product. All engine/solver updates were included on the same date as the first release.

Product	Release Date
OLI Studio/Analyzer	June 11, 2021
OLI Flowsheet: ESP / ESP FS¹	June 11, 2021
Chemistry Wizard	June 11, 2021
OLI Developer Edition	June 11, 2021
OLI Engine 11.x for Aspen Hysys (x64 bit)	June 11, 2021
OLI Engine 11.x for Aspen Plus (x64 bit)	June 11, 2021
OLI Engine 11.x for Petro-Sim (x32, x64 bit)	June 11, 2021 ²
OLI Engine 11.x for ProII	June 11, 2021
OLI Engine 11.x for Unisim Design	June 11, 2021
OLI License Manager³ (x32, x64 bit)	June 11, 2021
OLI Engine 11.x for gProms (x64 bit)	June 11, 2021

OLI Engine 11.x for IDEAS was not updated for this release

¹ The product ESP FS is for Asia releases. It is functionally identical to OLI Flowsheet: ESP.

² Version 11.0.1.x of the OLI Engine will only work with version 7.2 of Petro-Sim.

³ This product does not follow the same versioning scheme as the other products. The current version is 5.0.1.5

Databank Updates

Mixed-solvent electrolyte (MSE) databank

Special Note about the HF-Alkylation Database

In the V11 MSE databank (MSEPUB), OLI has incorporated all contents in ALKHF into MSEPUB. In this process, changes were made to the names of ASOs (Acid Soluble Oils), i.e., from ASO0, ASO1,...to ACIDSO0, ACIDSO1, etc. These changes are necessary due to conflicts of species names with variable names for activity coefficients (e.g. ASO2AQ and ASO3AQ for activity coefficients of SO2AQ and SO3AQ) - which caused problems in the comprehensive tests of V11 MSE databank by the Software Team. Because of these changes and updates to MSEPUB databank, OLI would like to make clarification as follows:

1. The ALKHF databank is no longer needed (**and should not be included - see 3 below**) in V11 for the HF alkylation modelling - due to the full incorporation of ALKHF into MSEPUB;
2. The ASO's can still be input as ASO0, ASO1,etc., as the MSEPUB keeps them as synonyms; but the chemistry model generated will be using the actual species names (e.g. ACIDSO0, ACIDSO1,...etc) in the related model files (such as dbs);
3. If ALKHF databank is included in V11, there will be conflicts of having the same ASOs with two different names (e.g., ASO0 (from ALKHF) and ACIDSO0 (from MSEPUB)) and different equilibrium equations (with different names), which causes issues as reported by Prodip.
4. ALKHF databank can (and should) still be used with V10 Studio and Flowsheet for HF alkylation modelling. The differences between V10 and V11 are in the use of different ASOs' names (i.e. ASO0 vs. ACIDSO0, which will not affect the results).

Acetic acid – acetate – hydrocarbon systems (revision)

Acetic acid – H₂O

Acetic acid – HCl

Na acetate – acetic acid

Li acetate – acetic acid

K acetate – acetic acid

Ca acetate – acetic acid

Acetic acid – CO₂

Acetic acid – acrylic acid

Acetic acid – methylisobutylketone (MIBK) – trioctylamine

Acetic acid – toluene – trioctylamine

Acetic acid – methanol

Acetic acid – ethanol

Acetic acid – hydrocarbons (methane, cyclohexane, benzene, ethylbenzene, o-xylene, p-xylene, m-xylene, isopropylbenzene)

Acids in carbon dioxide

H₂SO₄ – CO₂

HNO₃ – CO₂

Monoethylene glycol (MEG) systems (revision)

MEG – CO₂

MEG – CaSO₄

NaCl scaling chemistry (revision)

NaCl – H₂O pressure dependence

FeS scaling chemistry (revision)

FeS (mackinawite, pyrrhotite) – NaCl, HCl, NaHS,

Amine hydrochloride chemistry

Diethylenetriamine HCl

Monomethylethanolamine HCl

Dimethylethanolamine (DMEA) HCl (revision)

Morpholine HCl (revision)

3-methoxypropylamine HCl (revision)

Methyldiethanolamine (MDEA) HCl (revision)

N-Methylmorpholine HCl (revision)

Butylamine HCl (revision)

Dimethylisopropanolamine HCl (revision)

Isopropanolamine HCl (revision)

Methylamine HCl (revision)

Dimethylamine HCl (revision)

Trimethylamine HCl (revision)

Ethylamine HCl (revision)

Diethylamine HCl (revision)

Propylamine HCl (revision)

Sec-Butylamine HCl (revision)

Cyclohexylamine HCl (revision)

Ethylenediamine HCl (revision)

N-ethylmorpholine HCl (revision)

Ethanolamine HCl (revision)

Diethanolamine HCl (revision)

Diglycolamine HCl (revision)

Hydrogen fluoride - hydrocarbon chemistry

Hydrocarbon – HF – H₂O systems

Propene (C₃H₆)

Propane (C₃H₈)

Butene (C₄H₈)

Isobutene (i-C₄H₈)

Cis-2-butene (cis-2-C₄H₈)

Trans-2-butene (trans-2-C₄H₈)

Butane (C₄H₁₀)

Isobutane (i-C₄H₁₀)

Pentane (C₅H₁₂)

Isopentane (i-C₅H₁₂)

Neopentane (neo-C₅H₁₂)

Cis-2-hexene (cis-2-C₆H₁₂)

Hexane (C₆H₁₄)

2-Methylpentane (C₆H₁₄)

Cyclohexane (C₆H₁₂)

Methylcyclopentane (C₆H₁₂)

Heptane (C₇H₁₆)

2,3-dimethylpentane (C₇H₁₆)

Methylcyclohexane (C₇H₁₄)

Octane (C₈H₁₈)

2,4-dimethylhexane (C₈H₁₈)
Isooctane (C₈H₁₈)
Nonane (C₉H₂₀)
2,2,5-trimethylhexane (C₉H₂₀)
Decane (C₁₀H₂₂)
Undecane (C₁₁H₂₄)
Dodecane (C₁₂H₂₆)
Mixed hydrocarbon – HF – H₂O systems
Isobutane – n-octane
Isobutane – isooctane
Isobutane – alkylate
n-butane – isooctane
n-butane - alkylate

Acid soluble oils: ASO0, ASO1, ASO2, ASO3, ASO4, ASO5
Acid soluble oils – H₂O
Acid soluble oils - HF

Additional hydrocarbon chemistry

2-methylpentane – H₂O
2,3-dimethylpentane – H₂O
2,4-dimethylhexane – H₂O
2,2,5-trimethylhexane – H₂O

Mercury chemistry

Hg₂Cl₂
Hg₂Cl₂ – HCl, NaCl, KCl
HgCl₂ (revision)
HgCl₂ – HCl
Hg₂F₂
HgF₂
HgF₂ – HF – HgO – HgOHF

Cadmium chemistry

Cd(OH)₂ as a function of pH
CdO as a function of pH
CdF₂
CdF₂ – NH₄F

Osmotic pressure prediction

Improved algorithm for calculating the partial molar volume of water for predicting osmotic pressure

Phosphonate scaling inhibitors

HEDP (1-hydroxyethylidene-1,1-diphosphonic acid or etidronic acid)
HEDP complexes with Ca, Sr, Ba
NTMP/ATMP (Nitrolo tri (methylene phosphonic) acid)
NTMP/ATMP complexes with Ca, Sr, Ba

Zinc chemistry

Zn(OH)₂, ZnO as a function of pH (revision)
ZnO – NaOH
ZnO - NaCl

Lithium phosphate chemistry

Li₃PO₄, LiH₂PO₄
Li₂O – P₂O₅ – H₂O

Cobalt chemistry

CoHPO₄
Co₃(PO₄)₂, Co₃(PO₄)₂·8H₂O
LiCoO₂

Mixed sulfates of Li and transition metals

NiSO₄ – CoSO₄
NiSO₄ – MnSO₄
Li₂SO₄ – MnSO₄

Gluconates of transition and alkaline earth metals

Co gluconate
Fe(II) gluconate
Fe(III) gluconate
Mn gluconate
Mg gluconate
Ca gluconateRare

Rare earth hydroxides

La(OH)₃
Ce(OH)₃
Pr(OH)₃
Nd(OH)₃
Sm(OH)₃
Eu(OH)₃
Gd(OH)₃
Tb(OH)₃
Dy(OH)₃
Ho(OH)₃
Er(OH)₃
Tm(OH)₃
Yb(OH)₃
Lu(OH)₃
Y(OH)₃

Rare earth acetates

La acetate
Ce acetate
Nd acetate

Rare earth phosphates

GdPO₄, GdPO₄·0.667H₂O, GdPO₄·2H₂O
NdPO₄
NdPO₄ – H₂SO₄ – H₃PO₄
NdPO₄ – CaSO₄ – H₃PO₄

Rare earth citrates

Nd citrate as a function of pH
Nd citrate – citric acid - NaOH

Rare earth tartrates

La tartrate as a function of pH
La tartrate – tartaric acid
Nd tartrate as a function of pH

Rare earth sulfates in alcohols

Nd₂(SO₄)₃ – ethanol – H₂O
Dy₂(SO₄)₃ – ethanol – H₂O

Calcium sulfate chemistry

CaSO₄ – H₃PO₄

Fluoride chemistry

MgF₂ (revision)

MgF₂ – NaF including NaMgF₃

Alpha-methylstyrene chemistry

Alpha-methylstyrene – H₂O

Alpha-methylstyrene – acetone – H₂O

Alpha-methylstyrene – phenol – H₂O

Alpha-methylstyrene – H₂SO₄ – H₂O

TRIS chemistry

TRIS (Tris(hydroxymethyl)aminomethane) – H₂O

TRIS (Tris(hydroxymethyl)aminomethane) hydrochloride – H₂O

Additional urea chemistry

Urea - ethanol – H₂O

Sodium salts revisions

Na₃PO₄·12H₂O – additional solid phase

NaOH·Na₂SO₄·NaCl triple solid phase

Mixed acids

H₃PO₄ – H₂SO₄

MSE-SRK databank

Density improvements

CO₂ – H₂O

Elemental sulfur

S – H₂O

Acetic acid – acetate – hydrocarbon systems (revision)

Acetic acid – H₂O – hydrocarbons (hexane, cyclohexane, benzene)

Aqueous corrosion kinetics databank

Alloy 13%Cr – revision

Corrosive environments:

NaCl

Seawater – O₂

H₂S – NaCl

CO₂ – NaCl

CO₂ – H₂S – NaCl

KOH

NaOH – NaClO₃

Acetic acid

CO₂ – H₂S – NaCl – acetic acid

H₂SO₄

HCl

HNO₃
H₃PO₄
HF – HCl
FeCl₃ – FeCl₂ – NaCl – HCl
CuCl₂ – CuCl – NaCl – HCl
CuSO₄ – H₂SO₄
KCl
Completion fluid – CO₂
CaCl₂ – CO₂
Butyric acid
K₂CO₃
HCl – citric acid – methanol
Sour gas well simulation

Repassivation potential parameters:

Cl⁻
H₂S
SO₄²⁻
NO₃⁻
OH⁻
VO₄³⁻
MoO₄²⁻

Alloy 254SMO – revision

Corrosive environments:

NaCl
Seawater – O₂
Cl₂
Acetic acid
H₂SO₄
HCl
HClO
H₃PO₄ – NaCl
FeCl₃ – FeCl₂ – NaCl – HCl
CuCl₂ – CuCl – NaCl – HCl
H₂SO₄ – HCl – FeCl₃ – CuCl₂

Repassivation potential parameters:

Cl⁻
H₂S
SO₄²⁻
NO₃⁻
OH⁻
VO₄³⁻
MoO₄²⁻
S₂O₃

Alloy 316 – revision

Corrosive environments:

NaCl
Seawater – O₂
MgCl₂ – NaCl
NaCl – O₂
NH₄F
AlCl₃
Na₂SO₄
Na₂SO₄ – O₂
MgCl₂ – NaCl – CaCl₂

NaHCO₃ – NaCl
Na₂SO₄ – NaCl – NaAc – O₂
CaCl₂
BaCl₂ – O₂
KCl – O₂
NaBr
Cl₂
H₂S – NaCl
CO₂ – NaCl
CO₂ – H₂S – NaCl
NaCl – acetic acid – H₂S – CO₂
NaOH
NaOH – NaCl
Formic acid
Acetic acid
Formic acid - acetic acid
Boric acid
H₂SO₄
HCl
HNO₃
H₃PO₄
HCl – H₂SO₄
HCl – HNO₃
HF
H₂SO₄ – HNO₃
HCl – H₃PO₄
HNO₃ – H₃PO₄
H₂SO₄ – HF
HF – HNO₃
HF – H₃PO₄
HCl – HF
HBr – H₃PO₄
HBr – acetic acid
H₂SiF₆ – H₃PO₄ – H₂SO₄
HF – H₃PO₄ – H₂SO₄
HCl – H₃PO₄ – H₂SO₄ – H₂SiF₆ – HF
HF – HNO₃ – H₂SO₄ – HCl
CuCl₂
FeCl₃
FeCl₃ – FeCl₂ – NaCl – HCl
CuCl₂ – CuCl – NaCl – HCl
H₂SO₄ – Fe₂(SO₄)₃
H₂SO₄ – Fe₂(SO₄)₃ – NaCl
H₂SO₄ – FeSO₄
H₂SO₄ – CuSO₄
NaF – H₃PO₄
NaF – H₃PO₄ – H₂SO₄
H₂SO₄ – NaCl
H₂SO₄ – Na₂SiF₆
H₂SO₄ – CuCl₂
FeCl₃ – HCl
HCl – KI
CaSO₄ – H₃PO₄ – NaCl – NaF
HF – NaCl – O₂
HNO₃ – NaNO₃ – Cu(NO₃)₂ – NaF
NaCl – H₃PO₄ – O₂
HNO₃ – NaCl
Na₂SO₄ – NaCl – FeCl₃ – Na₃PO₄ – O₂
NaCl – Na₂CO₃ – NaOH
NaCl – CaCl₂ – CaSO₄ – O₂
KCl – NaCl – O₂

NaNO₃
CaCl₂ – KCl – KOH – K₂SO₄ – K₂CO₃
NaClO₃ – NaOH
H₂SO₄ – NaOH – SO₂
NaOH – NH₄OH – KOH
NaOCl – NaCl – NaOH
NH₄HS
NH₄HS – NH₃
NaCl – KCl – CaCl₂ – NaHCO₃ – Na₂HPO₄ · 2H₂O – KH₂PO₄ – MgSO₄ · 7H₂O – MgCl₂ · 6H₂O – Glucose

Repassivation potential parameters:

Cl⁻
Br⁻
H₂S
SO₄²⁻
NO₃⁻
OH⁻
VO₄³⁻
MoO₄²⁻
C₂H₃O₂⁻¹ (acetate ion)
NO₂⁻
S₂O₃

Carbon steel – revision

Corrosive environments:

I⁻¹

OLI Databook – Bugs & Features

Bug ID	Problem	Resolution
2748	When exporting a databank from the OLI Databook the fields in the A01 file do not line up with the data correctly ex: ORG shows Databank Name, ASID show ORG etc	Code was using column numbers with select * (all fields) so column index should be looked up by field name and not assumed. Updated code for A01 file to use a lookup. We should plan to update the balance of the A file exports to fix this. Also updated output of doubles to use a Fortran routine to match the output from ESP 9.6 databook. The only issue we are seeing now is ESP runs some fields through a double to float conversion which slightly changes the numbers. Talking with Arjun we should not do this to allow numbers to be more precise
3355	OLI Databook Make it easy to find the OLI Databook's	The system databases are now pinned so they are easy to find.
3356	A user requested that the search criteria be saved so that she can view the available list without all the duplications. Her preference is MATC. However, others may want formulas or synonyms. Therefore, save is the preferred improvement vs. fixed criteria that the person needs to change regularly	Updated code to support the options being saved (in the registry). Also updated initial startup code to stack the databanks and search windows for ease of use. Also update the search of species/interactions/mat codes to show the number of entries found.
3476	Chemical formula for GdO(OH) is misspelled as DdO(OH)	Resolved by TPM team.
3612	OLI Databook: Validate empirical formulas result in correct molecular weight on import	Updated code for saveGeneral to compute the molecular weight of the empirical formula and then compares it to the supplied molecular weight if they are not within a small tolerance the formula is wrong. If there is an error a message box will be displayed in databook, while converting an entire databank a warning file will be written into the same folder as the final databank.
3632	Databook: EXT-DENM is expected as DMD11-DMD15, currently this is BMD11-BMD15	Variables aligned properly
3730	The A04 file is exported in order entered not sequential order new synonyms are at the end rather than grouped with others synonyms for the same tag	Update the export code for A04 file to properly sort the exported data in tag/level order.
3810	When importing ASCII files if the version number in the A01 file is incorrectly formatted (ex: 10.) the converter would crash	Code placed to trap out this situation.
3821	When importing ASCII files if the version number in the A01 file is incorrectly formatted (ex: 10.) the converter would crash	Updated selected import code to use new import routines to solve the issue and ensure they are consistent.
3841	Update issues with Databook export of AFiles 1) Not V- on version number in A01 file 2) Date should be 00-mon-yy not 00-mon-yyyy 3) Aqueous record R_UQ/Q_UQ/RADI was output with trailing. for integer values 4) Dependent ref TRAN & QUAL fields do not match original	Updated SQLiteDataProvider and Databook to handle #1 correctly Update SQLiteDataProvider to fix 2,3,4

Engine/Solver - Features

Bug ID	AREA	Problem	Resolution
3534	Thermo	MSESRK Density Improvements - finalize + testing	finalized engine changes + conducted additional testing of cases. Continuation of OS-3447

Engine/Solver - Bugs

Bug ID	AREA	Problem	Resolution
3401	Thermo	Partial molar volume improvement for osmotic pressure calculations additional testing and cleanup	Tested, cleaned up and committed partial molar volume improvement for osmotic pressure calculations.
3640	Solver	OLI Engine is incorrectly predicting VLE/LLE for H2SO4/CO2	V9.6.2 predicts VLE until ~64 atm and suddenly change to LLE. V9.6.3 and after start predicting VLE below 64 atm but the predictions are not continuous. A known issue of the Engine is it tends to remove phases when it has difficulties to converge. Several bugs related

			<p>to Engine failing to predict L2 phase have been worked on for V9.6.3. Several “helpers” have been added to the solver to find the L2 phase before remove the phase equations. This helps the Engine to find the L2 phase at lower pressures for this case but still failed at some points. A few additional fixes are added to assist Engine to smoothly predict the L2 phase appearance.</p> <p>However, the L2 appearance at pressures lower than 64 atm is not physically reasonable according to Andre and Ron. But those are mathematically correct solutions so the Engine cannot avoid finding them. Ron found a workaround to start with an initial inflow ratio that avoids the LLE envelope.</p>
3886	Corrosion Solver	Update Rates valid information in Databank	Updated SQL file CorrosionRatesInfo.txt to included latest flags
4006	Solver	Crash in studio running MSE	Additional checks for pointer were added.
4061	Solver	Units for “Hardness” as “CaCO3 mg/L” were in error	Corrected the units per USGS documents.
4147	Solver	A case files fails with a solver error. If the flow units are in g/hr, then the calculation works fine. if instead it is changed to volume, then the calculation fails with the error shown. The problem disappears if a very small amount of H2O is added	A case files fails with a solver error. If the flow units are in g/hr, then the calculation works fine. if instead it is changed to volume, then the calculation fails with the error shown. The problem disappears if a very small amount of H2O is added. Added check to prevent a crash. The case will now catch the error. The volume target may still fail as this type of gas only calculation does not go through the solver without water, hence, volume target can not be achieved. A controller may be used alternatively. amount of H2O is added

OLI Studio/Analyzer - Features

Bug ID	AREA	Problem	Resolution
155	Plotting	Need to reorder variable columns in the View Data for exporting	Reorder implemented.
645	Plotting	Units do not change on plot after changing the via Units Manager	Plot is now updated after units changed.
919	Units Manager	There is a need for mmol/L in units	Added

OLI Studio/Analyzer - Bugs

Bug ID	AREA	Problem	Resolution
---	Copy and paste	Copy as stream from OLI Flowsheet to OLI Studio that has no flow for H2O, the paste may have invalid values for water (typically a value related to 55.5082 gmoles)	No resolution for V11, a known issue to be fixed in a later version.
392	Stream Analyzer	Autoclave Calculation does not report some thermodynamic values	In the autoclave calculation, the array "iprop" controlling the output of thermo properties is first defined and allocated correctly in the subroutine "eqautoclave". However, when it was passed to a subroutine "user_model5", it was deallocated and re-allocated with different size and then initialize with different values. This affects the properties being reported. Both deallocation and subsequent allocation of "iprop" are unnecessary in "user_model5", and thus being removed.
564	Stream Analyzer	Copying a mixer removed the inlet stream results in the source mixer	When a regular survey is copied all survey points got temporarily removed, and then restored after copy. In mixer there may be flash calcObjects which are not survey points. When these flashes are temporarily removed their reference count drop to 0 and self destructed. To avoid that we temporarily add these flashes to the survey point array, and later remove them from the array. Thus the flashes will not get destructed after survey point array restored. The other issue is when these flashes were temporarily removed the m_keyScratchPadCalc were set to -1, so we have to set them back the the valid key values during the restore.

647	Corrosion Analyzer	Settings for a polarization curve are not saved after a file save.	The bug is due to always use the first line's color for all the lines in a given sub group. However, once user has changed individual colors in the sub group we should honor user selection.
746	Stream Analyzer	Copying data from an external source into a brine analysis did not work if the source data had commas (instead of decimals) in the data.	Updated number parser to support comma's looking for correct number of digits between commas and proper placement of comma's in the number. Incorrect placement and the number will not be parsed.
1673	Stream Analyzer	Calculation with REDOX enabled used the "Save As Stream" function but used the original inflows and not the inflows based on the reduced or oxidized species.	New inflows are used in the "Save As Stream" function based on the calculated redox results and not the original inflows.
1744	Corrosion Analyzer	Pourbaix Diagram has incorrect contact surface when is copied and pasted	Updated code to properly copy contact surfaces when they are a component in the databanks. Copying a diagram that uses an allow still does not work, this is a larger issue and a new story will need to be opened to address this.
2244	Solver	Incorrectly calculated phase behavior (100 % all vapor when 2 phases are expected) in a survey	Vapor-organic initializer for the solver was updated.
2256	ScaleChem	A saturation calculation incorrectly calculated the neutral H2S concentration.	The internal tables used to fill in the value in the report used an incorrect molecular weight.
2261	Framework	A stream copied from OLI Flowsheet in "Day" units had its flowrate multiplied by 24 in OLI Studio, also in Day Units	Conversion error fixed.
2271	Solver	<p>Modified critical parameters of pseudocomponents in MSE-SRK for better match. EOS solver cannot solve properly for the phases, as can be seen from the jump to 10 or more. (Insert: before modification)</p> <p>The moles of the vapour phase suddenly jump from 0 to 10 when the solver finds a vapour phase (i.e. in the 2-phase region). However, the phase transition for a mixture of components should be gradual. Starting with a tiny bit of vapour at the phase boundary, which gently increases as P,T moves further into the V+L region. See for comparison the graph for the un-modified pseudo-components.</p> <p>I interpret this as an instability in the solver when trying to solve for a condition close to the phase boundary, which seems to be exacerbated when I have modified pseudo-component properties. The latter action may destroy some correlations that normally apply, but I don't know if this is triggering the behavior.</p>	This MSE-SRK case has a hydrocarbon-only stream, and cannot predict vapor appearance at increasing temperature. The Engine uses the successive substitution method with the Rachford-Rice algorithm to solve SRK VLE, however, has a convergence issue. The vapor pseudo-root extrapolation on a supercritical fluid in the SRK subroutine causes the discontinuity and fluctuation in convergence. By changing the SRK subroutine to use identical vapor and liquid roots for supercritical fluid, and tuning the Rachford-Rice algorithm, the solver is able to continuously predict the vapor phase appearance.
2332	Interface	Distillation (assay) data entry does not have the units for the type	Assay data entry now displays the distilled value in volume or mass percent as indicated by the method.
2360	Corrosion	Inflow species values for a corrosion rate calculation were locked when switching to MSE-SRK	Corrosion rates are not value for MSE-SRK. Error was trapped.
2557	Framework	A stream with redox enabled copied from OLI Flowsheet to OLI Studio did not keep the redox species enabled	User interface updated to honor the selections made in the host program.
2593	Corrosion	After using the "Set pH" calculation and then switching back to "Isothermal" the titrant selection is still displayed.	Updated code to track which calctype is set for which survey var/covar is selected. This is used to reset the calctype when the user changes the selection. Also exposed calculation type in the grid.
2714	ScaleChem	CRASH: Program runs and exits without any warning	Found issues with code where crashes could occur and fixed
2719	Framework	Tool tips are missing for contour plots.	This is implemented. We created a OLIToolTipCtrl class that override some of MFCToolTipCtrl functions and fixed the problem that tool tip won't display if control is disabled. Now OLIToolTipCtrl will display tool tip even if control is disabled. The OLIToolTipCtrl is now in the OLIControls project.
2969	Solver	A case with the hardness calculation enabled failed on every other point in MSE. If the restart vector was enabled, then all points converged.	A problem with the call for the hardness calculation was identified and corrected.
2980	Solver	A case with surface complexation enabled (via the database) crashed on the report. The calculation is converged.	When user uses mass fraction or mole fraction, in solid phase we provide total mass or total moles from solid phase. This will end up getting 0 if all solids are in the surface species. To fix this we must know if user is asking for total from solid phase or surface species and provide the correct totals. This is fixed for phase table, true species table and MBG table. Also, the output grid for

			these sections is display correct values. We have also tested concentration units and they are not affected.
2988	ScaleChem	The brine composition table displays the values in mg/L regardless of unit selection. The column headings do reflect the user choice however.	This is fixed. The old code only allowed mass or mass concentration units to be used for brine table. This is not true anymore as Studio now allows any unit user selected to display the numbers.
2994	Stream Analyzer	If a calculation in volume units fails to converge then the parent stream becomes locked, and thus no new calculations can be added	If converge volume failed the parent stream did not have a chance to get unlocked. This is fixed by unlocking the parent.
2996	ScaleChem	Saturator Calculation failed with heavy hydrocarbons and solid phases	The gas stream is methane dominant, and the condition is in its critical region, i.e. $T > 100$ C and $P > 200$ atm. The solver is having an issue to verify as vapor or organic phases since both phase checks passed. The solid inclusion calculation is inconsistent in the organic phase check and not able to converge. The fix is able to make all these three bugs converge.
3274	ScaleChem	A reconciled brine is double-accounting the reconciled ion in the brine report	The ReconcileCalc under brine set result calc to true so it does not have the original input values. When the report requested the inflow values it provided the output values. This is now fixed by using the brine as the input stream to provide the correct input values.
3377	ScaleChem	A solid is missing in the selection list because the first point in the contour plot is zero. If Hide-Zero species is unchecked, then the solid appears.	Fixed issue with the logic when the All-solids option was selected and a solids with a 0 at the first survey point was not showing. Now correctly shows up when hide 0 species is enabled.
3378	ScaleChem	In the plot variable selection list, the Standard solid v. Expanded/All solid filter always reverts back to standard if leaving and returning to the dialog. Once selected it should stay selected.	This is fixed by adding a CSting member in PlotInformation to keep track user selection.
3485	Stream Analyzer	The variable TDS, when enabled, appears in the output tab but not in the report	This is fixed. Now TDS is in reports of Single Point, Survey, and ScaleChem calculations.
3606	Stream Analyzer	The mixer was set up as an isothermal calculation at 23.6 C, yet the flash calculations are being performed at 180C. TDS had been enabled	The mixer case takes in streams in concentration units and need to do volume converge when creating flashes for the streams in the preCalculate() call. During preCalculate() EQTDS was called which changed the GVEC, and later on when the calculate() is called the modified GVEC was used (temperature is 180C). To fix this EQTDS should not be called during preCalculate(). Chris suggested during preCalculate() we should turn off all calc options. The calc options are only for the real calculate() call. After turning off all calc options during preCalculate() the issue is resolved.
3713	ScaleChem	A brine that was copied and then manually updated (as opposed to a deletion) did not reconcile correctly.	This is a bug due to fixing OS-2075: Brine reconciliation had different calculated pH's if calculate alkalinity is checkedDONE in which H2O input value got corrupted. We fixed it by forcing it to be 55.508 mol during post serialization. However this is only true if the brine is in mass concentration units. If brine is in mass fraction units such as ppm the H2O input value can vary. Forcing it to 55.508 mol in post serialization will cause errors. The paste action does end up calling post serialization. To fix it we only make sure H2O is 55.508 when the units are mass concentration. Otherwise don't change H2O value in brine.
3714	Stream Analyzer	Gibbs Free Energy value are reported as zero in report. They are not zero	The Gibbs Free Energy value of each species is non-zero. However, if the amount of a species is 0 in a stream, its Gibbs free energy values are not reported. To solve this, I first removed the explicit check of species amount to allow the Gibbs free energy each species to be reported. Then, I also removed the check of the species amount in a particular phase.
3736	Stream Analyzer	Arsenic is missing from possible redox subsystems.	The redox import table that creates the Redox Information table was missing these systems for MSE. The following systems were missing or had incorrect information Arsenic, Bromine, Chlorine, Curium, Mercury, Phosphorus
3788	Stream Analyzer	Enabling the TDS calculation in a mixer give a different titration curve.	This is the same issue as OS-3606: Bad Isothermal Mixer CalculationDONE the TDS calculation is being applied to

			the inbound streams and changing the temp to 180. This causes the rest of the calculation to be incorrect.
3890	Solver	Using a species with a dimer (acetic acid dimer for example) and MSE-SRK results in a doubling of the mass for the inflow.	The bug is confirmed when the flashed result is in organic-only condition. When the solver copies flash results of the organic phase, it didn't overwrite log(x) variable if x is 0 (to avoid log(0) error). However, it should be overwritten by a very negative number, i.e. -300. After the fix, the organic composition is reasonable without double counting of acetic acid dimer/monomer.
3917	Stream Analyzer	Case created in V10 in mg/L units fails to converge.	Added code to restore missing species and make sure water is not zero in concentration unit.
3931	Corrosion Analyzer	In a case with redox, a sub-system was missing in the drop-down menu from the "RE" button.	This is fixed by making sure the search list obtained valid redox subsystem data. The bug is due to cached redox subsystem is invalid and the name is empty. If this happens we should discard the cached data and look for valid data in the databanks. See attached fixed image.
4037	Stream Analyzer	OLI Studio summary view not showing consistently	Found stray code that prevented this from appearing.
4093	Units	Hardness values are wrong and with wrong units	Changed unit for hardness to 'mg/l as CaCO3' in OLIValues Updated OLIStudio and OLIFlowsheet to correctly set the new unit on reload of case and to remove the hardness value from old cases as it is incorrect.
4115	Solver	CRASH: case that is incorrectly set up quits with no error. This is a set pH calculation with zero water. It is an improperly set up case. The program quits without any kind of error message or stack dump. We should prevent these sudden crashes and warn the user that the case is invalid.	Added check to prevent the crash. Case is invalid but now will catch the failure without crashing.

OLI Flowsheet: ESP / ESP-FS - Features

Bug ID	AREA	Problem	Resolution
3597	Solver	isochoric calculation improvement for OLI Flowsheet	an iterative isochoric calculation method has been implemented [only for flowsheet]. Set as alternative method when regular method fails.

OLI Flowsheet: ESP / ESP-FS - Bug

Bug ID	AREA	Problem	Resolution
612	Import	Copy calculated stream from Studio (redox enabled) and paste-update in Flowsheet, does not carry information for redox	<p>When copying a stream from Studio to Flowsheet the behaviors are explained in the following user options:</p> <p>The stream will be pasted to a new stream in Flowsheet using the default Chemistry Model. The pasted stream will be identical to the source stream and the default Chemistry Model will be merged by source stream's chemistry. Redox will honor the source stream's redox if there is overlap.</p> <p>The stream will be pasted to a new stream in Flowsheet in a new sub Chemistry Model. The pasted stream will be identical to the source stream and it uses a newly created Chemistry Model (sub Chemistry Model) in Flowsheet. The default Chemistry Model is not touched.</p> <p>The stream will be pasted to an existing stream in Flowsheet using the default Chemistry Model. The existing stream will be merged with the pasted stream and the default Chemistry Model will be merged by source stream's chemistry. Redox will honor the source stream's redox if there is overlap.</p> <p>The stream will be pasted to an existing stream in Flowsheet in a new sub Chemistry Model. The existing Flowsheet stream will be replaced by the pasted stream and it uses the newly created Chemistry Model (sub Chemistry Model) in Flowsheet. The default Chemistry Model is not touched.</p>

615	Solver	Column heat exchanger duty is not correctly transferred using a heat transfer block. If the exchanger heat duty is varied for a target, the final converged heat duty is not correctly transferred using a heat transfer block. Only the initial estimate is transferred. In the attached case, column initial estimate 1.0 is transferred but not the final converged value of 1.20791e5	Added/corrected logic to transfer heat. There are a few combinations possible such as Transfer in, transfer out. Testers need to test all possible combinations.
1571	Solver	Scaling tendency for Fe3O4 is non-zero when one of the components has a zero flowrate	The bug is in the settler block where scaling tendency indices are messed up. It was double converted to the master model index. After the fix, the output shows the correct scaling index for species.
1584	User Interface	Large cases take a long time to export a report	Tested again and found high latency across VPN connection, updated code to write to a local temporary file and then copy to the network which improved the speed.
2134	User Interface	Could not entirely delete a virtual stream.	When deleting a virtual stream, we must initiate the delete from the parent portal so both the portal and the virtual stream will be deleted.
2190	User Interface	Energy Transfer Block heat duty not displayed in block call-out	User Interface updated.
2391	User Interface	The interaction matrix is not available (is available in OLI Studio)	Added menu item View Interaction Matrix (csv) File to launch user app associated with csv file.
2501	User Interface	The navigator panel does not sort alphabetically when that option is selected.	This option is now properly honored.
2540	User Interface	A stream appears on the PFD but not in the navigator. Cannot delete the stream.	Fixed issue that caused stray/defunct external inlets to a block due to a failed stream reconfigure operation. Added code to handle phantom visual streams from old, saved documents.
2649	Reports	Streams with no liquid phase should not have a zero-pH reported	Report updated to show a blank field in such cases.
2651	Reports	Multi-stream report, when created stream-by-stream, in the order the user wanted is reset when leaving the report	Fixed error in program logic when a stream is added to an empty column of the report.
2800	Framework	A file with reaction kinetics failed to open, program crashed.	There were invalid kinetic parameters in 4 equations. Updated code to ignore these parameters rather than abort the file open.
3004	User Interface	Could not delete a virtual stream (see 2134)	Fixed issue that caused stray/defunct external inlets to a block due to a failed stream reconfigure operation
3132	User Interface	Clearing results should result in a file changed state – the “*” that appears in the title bar.	Added the changed state when clearing results.
3180	User Interface	Pausing a calculation does not allow the user to edit the block that was paused.	Editing now enabled.
3260	User Interface	Multistage block callout did not display the correct stage number when selected	The interface was correct but confusing. It was cleaned up to show how to pick the correct stage.
3305	Solver	Sensitivity plot results do not match analogous case in OLI Studio	This was resolved with using proper restart values.
3308	Solver	Start/Stop/Restart does not work with virtual streams	Updated calculation order fixed when using virtual streams.
3309	User Interface	Enter key does not advance the cursor in sensitivity panels.	Tab/Enter key updated.
3338	User Interface	Hide-zero function does not work in call-outs	The hide-zero functionality was now properly implemented.
3348	User Interface	Cannot use the restart functions for a controller which is connected to an internally manipulated stream	The restart selector algorithm was expanded to search for internally manipulated streams.
3352	Solver	Inconsistent phase predictions at high pressures	The vapor phase starts to appear at higher pressures, and the vapor density shows the phase is from SRK liquid root. When SRK cannot solve vapor root at such high pressures, an extrapolation method is supposed to be used to report high fugacity coefficients. However, this case the SRK subroutine also tried the analytical solution of SRK root and incorrectly used the liquid root. A stability check is added following the new engine implementation and fixed the bug.
3379	User Interface	Feedback controller appears to converge to an incorrect set point with multiple models	The controller should get the master model. Since old ESP does not write block specific chemistry model for the controller (feedback, feedforward, and sensitivity too), the engine makes sure these blocks are run on master model. Flowsheet ties model name to some of these blocks and that creates confusion to the engine. In this case, the controller is tied with a sub-model and that causes the mentioned issue.

3381	User Interface	Pressing Enter on the spec value in a multi-stage column spec/control closes the dialog. It should move to the next available value or button.	Tab/Enter key updated.
3383	User Interface	Earlier versions allowed the direct enter of a stream name in the block panel.	This has been re-enabled.
3402	Solver	Partial molar volume improvement for osmotic pressure calculations - RO membrane side	Some code in RO membrane side needs to be replaced and tested to be consistent with partial molar volume improvement.
3452	Chemistry Model	When MSE-SRK is selected but a non-suitable species is entered a warning should be issued	Database updated and warnings now displayed.
3471	User Interface	A stoichiometric reactor fails without a reliable error message if there are more than 7 reactions.	Added an error flag to inform the user that there can be no more than 7 stoichiometric reactions.
3475	User Interface	A feature that worked in earlier versions, copy-paste a stream call-out fails.	A lot of code modification has happened between v10 and v11. After a thorough code mapping the bug is pinpointed at a few lines of change. Also added is the serialization of m_bUseSrcUnit which makes sure units are honored after paste.
3477	User Interface – Report	A case file has three water analysis sampl. The report for the samples always uses the last sample regardless of the selection.	This is fixed. WaterAnalysisResultsGrid class has a member m_linkWA which contains the valid water analysis object. It was first set in the setReport() function and never changes. When user click the dropdown list to select another water analysis object, m_linkWA is still using the old object. The fix is to update m_linkWA to contain the newly selected water analysis object. This is done in WaterAnalysisResultsGrid::getObject(). The other bug fixed is that water analysis report always shows “Stream” in the header row rather than the actual name of the water analysis object. This is fixed by overriding getTitle() in the WaterAnalysisResultsGrid class to get the correct name.
3480	Solver	A case with isochoric mixers fails but works with standard mixers.	Improved isochoric calculations, see OS-3597
3587	User Interface	Calculation sub-type was not saved if block was configured to be an isochoric calculation.	This was corrected
3588	User Interface	Adding salt rejections as ions to the RO block automatically adds the ions to inflow list	Check to see if species is already in the chemistry model. If not add it. Also check if ions are allowed. This is now fixed.
3592	User Interface	Restarted value for a heat exchanger was many orders of magnitude less than the original value.	Unit conversion error.
3631	User Interface	Heat Exchange set to vapor fraction in mass % units. The value calculated was in mole % units.	Mass % units are not supported. UI updated to accept only mole fraction or mole %
3642	Solver	Sensitivity study returns an error: Subscript out of range on warning vector - crashes engine	Subscripts fixed
3690	User Interface	Call-out for a compressor block was missing some parameters. Denoted with “?”	All parameters now properly populated
3712	User Interface	Direct flow manipulator does not update callout unless you close and reopen	If the block controlling the stream and the stream has the same name, the update does not occur. This confuses the block-order calculation and is now prohibited.
3755	User Interface	Callout for multistage column only reports a single heat exchanger. There are multiple heat exchangers	The interface was updated to make it more intuitive to select multiple exchangers.
3756	Import	Stream copied from OLI Studio (used water analysis) has much different mole flowrates in OLI Flowsheet	Since the source stream's inflows were in mass-by-vol conc. basis which are not supported in Flowsheet they were converted to mass basis. However, they were not being scale to the total stream amount (even though they would be automatically scaled during calc time as the total stream was correctly overridden). Modified stream paste logic to translate inflows in mass-by-vol or mole-by-vol conc. basis into mass-fraction/mole-fraction basis as it would be the closest substitution of the original inflow amounts.
3761	Solver	Direct flow manipulator versus traditional manipulator gives vastly different results - UI work for unique block and stream names where necessary	This is fixed by enforcing unique name for controller's target streams. It is generally allowed for a stream and a block to share a same name. However, for a controller's target stream the name must be unique. When opening an existing file this rule will be applied by checking all controller's target streams. Errors will be flagged if duplicate names are found. User must fix them before calculation can proceed. On the other hand, if a user tries to change the name of a stream or a block the same rule

			will be applied if the system has target stream of controllers. See OS-3712
3766	Solver	Zero enthalpy reported for liq2 only calculation	Fixed a new bug for not calculating enthalpy for Liq2 stream, and Corrected a previously fixed bug showing some extractor cases failing.
3779	User Interface	Add Electro neutralities information to in the definition tab to Water Analysis tool in Flowsheet	Added electroneutrality information and other calculated properties to the calculated results.
3780	User Interface	Feedback controller target should not allow connecting to block - stop in UI	The target object of a feedback controller (input to the controller) should be stream object only. The flowsheet view does allow user to drag and drop on a block object, which will cause an error. Now the code has prevented the drag and drop on a block object.
3781	User Interface	Clear results deletes the column restart data for flowsheet	Modified logic for the "clear results" action to not clear tower restart data.
3809	Chemistry Model	Ions entered in chemistry inflow grid are not displayed with the "erroneous inflow" formatting	In chemistry inflows entry grid added logic to identify ions that may have been entered by the user and use "erroneous" formatting to flag them. An error tooltip is also shown for inflows that are ions.
3880	User Interface	Turbine default efficiency is empty for old cases opening in V11, showing error - new cases not running too	Corrected handling of default values for the 3 different types of efficiencies in the Turbine's specification.
3935	User Interface Reports	The Case file uses ion exchange. The Adsorbed View table displays the materials on the surface species. The S-5 table contains the surface MBG and Combined MBG. The surface variables are zero. This is due to a units Manager issue. The Report 2 will contain the values that should be present.	This is fixed: (1) The MBG callout will display correct info now. (2) if there is no Aq phase and user wants to use concentration or molar concentration units, we will automatically change to mass fraction or molar fraction units to accommodate the stream with no Aq phase. Please see attached before and after images for Bug(1) and Bug(2).
3937	Solver	This case has two models Chemistry (default) and REDOX. One includes iron redox. If I set one block to REDOX, the block fails (much further upstream)	When a new chemistry model is introduced into the process, the structure of the saved restart vectors becomes inconsistent with the new vectors setup by the Engine. Added logic in start/stop/restart optimizations code to properly handle such situations.
3939	User Interface & Solver	Feedback controller and target heat duty in a heat exchanger are not using the same set of units.	Wrong blmain units were being retrieved when converting feedback controller's user-specified param bounds for writing to bin file.
4005	Solver	Disabled controllers in case file still report as having not converged.	Disabled controller reported to fail in error. Dispatch object update notification to ensure observers are notified when list of calc-errors is changed. Also skip reporting special feedback controller converge fail error if said controller is no longer enabled.
4050	User Interface	Flowsheet UI freezes when the definition pane for a block is visible while a simulation is running for a very large flowsheet.	The block's spec grid was updating constantly when any flowsheet object's calculation state changed during the simulation. This blocked other UI updates from happening and the UI appeared frozen until the calculation finished. Narrowed down the filter for events that triggered an update of BlockSpecGrid to prevent unneeded updates.
4078	Solver	Error in multiple models (to be fixed in a later version) crashed and did not exit gracefully	Error not properly trapped. Program does not crash but still persists in this version.
4093	Units	Hardness values are wrong and with wrong units	Changed unit for hardness to 'mg/l as CaCO3' in OLIValues Updated OLIStudio and OLIFlowsheet to correctly set the new unit on reload of case and to remove the hardness value from old cases as it is incorrect.
4103	Solver	Program hangs when using a water analysis object	Deadlock when water analysis calc is running. Keep cached copy of dependency chem model so that getChemistryModel() is not called unnecessarily. Calling getChemistryModel() may be computationally expensive if the source water analysis's chem model needs to be synchronized.
4131	User Interface	A controller parameter set was not found in the species list. This was an ion concentration in a 2 nd liquid phase in MSE-SRK which has no ions in the 2 nd liquid phase.	We are incorrectly writing CLIONAQO to the bin file it should be written as CLIONO
4142	User Interface	Calculation failed in converge volume. No Trace File	Updated code to enable trace file for the converge file calculation (if enabled in parent) and to show converge volume failed if that is the case.
4172	Import	Importing a case from ESP Original had a stream in mole fraction units. There was no water present in the stream. A mole fraction of 1 was added for water on import.	Error in stream composition after import: Fixed issue with import defaulting water to value of 1 when fractional units are specified and since the stream did not have an entry for 0 WATER the water remained at 1. Now explicitly check if water is included in the incoming stream and if

			not explicitly 0 the entry for water. Oddly, this worked inversion 9.6 but not in version 10
4185	Solver	Program crashed with an "Unexpected Way" message, suspect too many water samples	<p>Found issue, when setting units in a water analysis you can check the option to use these units for all new objects, this allows new flowsheet objects(Mixers...) to be concentration.</p> <p>This was found to be caused by concentration units being set for a mixer block which is invalid. It was tracked down to the units Manager allowing Water Analysis units to be set as the default for the document. Updated the Units manager to hide this option when it is a Water Analysis object.</p>

Alliance Products - Features

Bug ID	Program	Problem	Resolution
----	Petro-Sim	Installation of version 11 of the OLI Engine will not work with Petro-Sim v7.1 or earlier	This is a change enforced by KBC (publisher of Petro-Sim). The OLI Engine version 11.x will only work with v7.2 of Petro-Sim or later.
3983	Unisim Design	OLI Configuration tool did not understand about Unisim Design R40.1 and R490	XML file updated

Alliance Products - Bugs

Bug ID	Program	Problem	Resolution
3192	Petro-Sim	If you open up an existing OLI case in PetroSim and click the about button for OLI, prior to any other operation it will crash.	Added protection code around pointers used for the about dialog.
3741	Aspen Plus	Very different results when a zero flowrate species is included in the stream	Updated initializer to enable organic phase check. Initial tests ran OK. Need to run against release tests with new databanks.
3999	Petro-Sim	Redox not being honored in model creation in Petro-SIM using chemistryBuilder	This is a bug in the chemistry builder that need to rebuild the system as needed when specifying redox. It's working correctly after the fix.
4014	Petro-Sim	KBC chemistry builder redox subsystem incorrect for hydrogen/oxygen	<p>In current redox logic, H2O redox is shown when H2 or O2 is in inflow. It is redesigned that H2 will not bring in H2O redox, and O2 brings in redox options for ozone and H2O2.</p> <p>Additionally, redox subsystems like FEEL are selected by default based on databank information. This feature was not there.</p> <p>Also added function to get databank full name for display.</p>

OLI Chemistry Wizard (all products)

Bug ID	Program	Problem	Resolution
733	All	The Chemistry Wizard for assays should reflect the proper units. For example, the percent distilled for method D2887 should be weight percent and we report volume percent.	Resolved for OLI Studio OS-2332
3547	All	Read MOD files without making any changes to them. Honor DEFINE equations	If a cws file is created from a mod file containing user defined equations, these equations will be added to the cws file. When a new mod file is generated from the cws file, these equations will be written to the new mod file.
3645	Regression	CRASH: Chemistry Wizard for Regression crashes out without error message	Tracked down to an issue in dbAutoSQL.for, array for redoxSpecies was allocated to the number of species which is not large enough for this case. Changed to allocate number of species*4. Test and issue resolved.
3882	All	Chemistry wizard looks at the old options to determine if MSE and SRK are enabled. This needs to be updated.	Updated code to use databank info from DataProvider, and to stop looking at the license. DataProvider only returns licensed databanks.
3972	All	Chemistry wizard crashes in autogen with selecting default Redox for many species	This is caused by the same problem as OS-4015: Chemistry wizard brings in ammonia related species N2 inflow, redox turn on but no redox subsystems selected DONE that Engine tries to include all missing redox systems when the

			redox is turned on but no subsystems are selected. It is fixed together with the other Jira story.
4002	All	Crash when selecting default REDOX for Ti species	Arrays updated to accommodate the size needed.
4174	gProms	Incorrect error message when using a private database	All databases are now stored in My OLI Cases/Databanks. This check is no longer needed.

OLI Developer Edition - General

Bug ID	Program	Problem	Resolution
		There were no upgrades except for overall solver/engine upgrades	

OLI Framework - General

Bug ID	Program	Problem	Resolution
3074	Security	Add new options in the license to support Database Segmentation, Databook and Regress, Flowsheet Pkg Publishing, named user flag	This is being deployed to new customers of OLI
3896	Support	Update Contact support to Show Portal not Support email	Update screens to show the portal and wiki , not email and wiki
3968	Security	Licenses for third-part software was out of date	Updated the licenses and copyright data in the security tab.
4190	Security	Named Users does not work for OLI Flowsheet	Named Users log in used the wrong screen. Now updated.