

ICON8 trials programme

N=1485

ICON8

ICON8B

N=1170

Stage IC-IV EOC/PPC/FTC

Randomise 1:1:1

Arm 1
6 cycles

Arm 2
6 cycles

Arm 3
6 cycles

Arm 1 Carboplatin AUC 5 q3w
Paclitaxel 175mg/m² q3w

Arm 2 Carboplatin AUC 5 q3w
Paclitaxel 80mg/m² q1w

Arm 3 Carboplatin AUC 2 q1w
Paclitaxel 80mg/m² q1w

High-risk* stage III -IV EOC/PPC/FTC

Randomise 1:1:1

Arm B1
6 cycles

Arm B2
6 cycles

Arm B3
6 cycles

Maintenance bevacizumab
(18 Cycle Total)

6-weekly follow-up
until week 66
post
randomisation

Maintenance bevacizumab
(18 Cycle Total)

Arm B1 Carboplatin AUC 5 q3w
Paclitaxel 175mg/m² q3w
Bevacizumab 7.5 mg/kg q3w

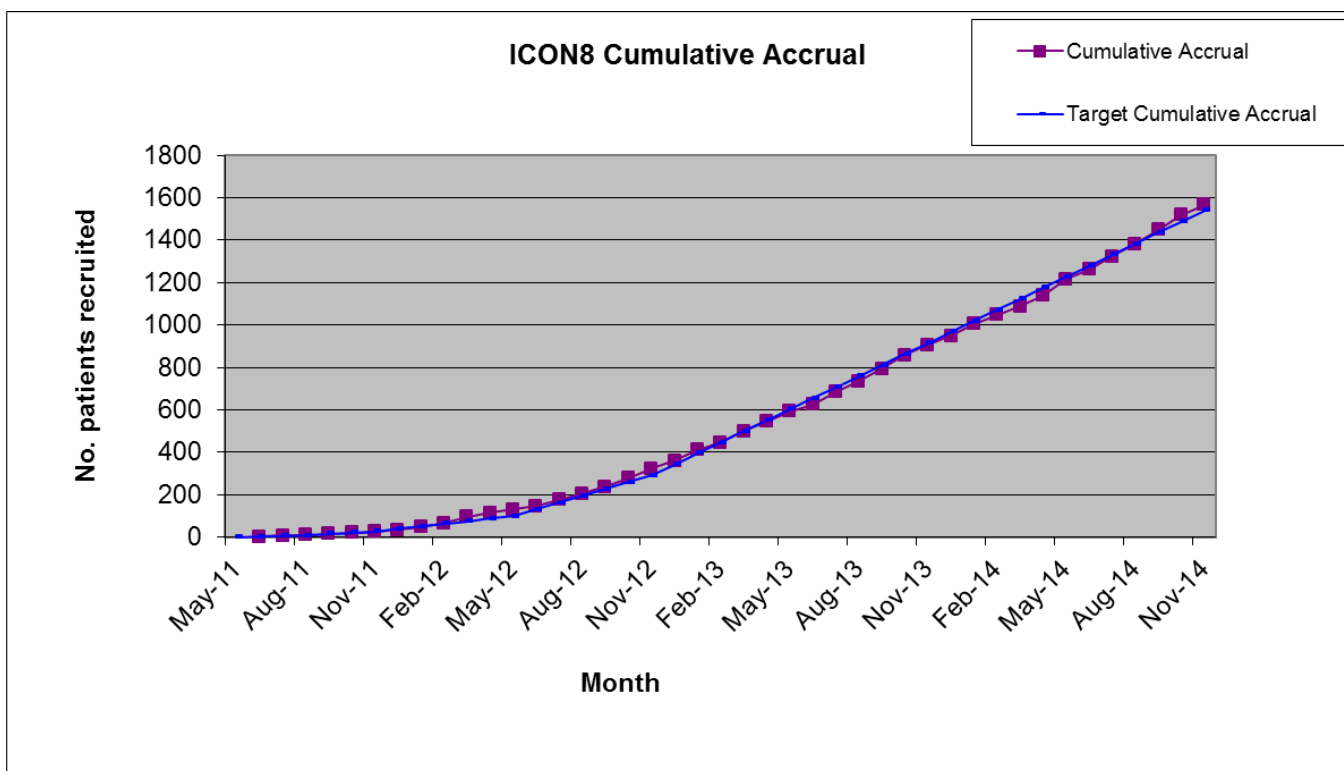
Arm B2 Carboplatin AUC 5 q3w
Paclitaxel 80mg/m² q1w

Arm B3 Carboplatin AUC 5 q3w
Paclitaxel 80mg/m² q1w
Bevacizumab 7.5 mg/kg q3w

NB. High-risk patients remain eligible for ICON8 so that patients with contra-indications to bevacizumab and those unable to access it are still able to enter the trial

High-risk defined as (1) FIGO (2013) stage IIIA1(ii), IIIA2 with positive retroperitoneal lymph nodes >1cm in diameter, stage IIIB or IIIC with >1cm residual disease following immediate primary surgery or planned to receive primary chemotherapy +/- delayed primary surgery and (2) FIGO (2013) stage IV

- Accrual began 06/06/2011 and ICON8 pathway closed to recruitment 28/11/2014



- Final recruitment figure = **1566**
- UK= 1397, ANZGOG= 70, GICOM= 43, KGOG= 32, ICORG= 24



ICON8 Outcome measures & analysis

Presentations:

- ESGO, October 2013 - oral poster presentation on stage IA analysis
- NCRI, November 2013 - poster on stage IA analysis; NCRI award for abstract
- ASCO, June 2014 – poster

- ❖ **Stage IA** showed that the weekly regimens were harder to deliver but total doses and dose intensity were increased. Uncomplicated grade 3/4 neutropenia was higher in Arms 2&3 but other toxicities were similar. Earlier use of GCSF was recommended following this analysis.
- ❖ **Stage IB** was reviewed by the IDMC in Nov-13. They considered the regimens safe and feasible for neo-adjuvant chemotherapy. DPS was not compromised in the weekly arms.
- ❖ **Stage 2** Activity Outcome measure: 9-month progression free survival rate in 1st 186 women randomised Completed Jan-14. Analysis reviewed by Independent Data Monitoring Committee, decision to continue all arms
- ❖ Anticipate **Progression Free survival analysis Q4 2016 & overall survival analysis Q4 2018**



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ICON8B

A study of bevacizumab and weekly dose-dense paclitaxel in ovarian cancer

Arm B1	Carboplatin AUC 5	q3w
	Paclitaxel 175mg/m ²	q3w
	Bevacizumab 7.5mg/kg	q3w
Arm B2	Carboplatin AUC 5	q3w
	Paclitaxel 80mg/m ²	q1w
Arm B3	Carboplatin AUC 5	q3w
	Paclitaxel 80mg/m ²	q1w
	Bevacizumab 7.5mg/kg	q3w

Aim to recruit 1170 participants over 4 years in 80+ sites across the UK and Ireland

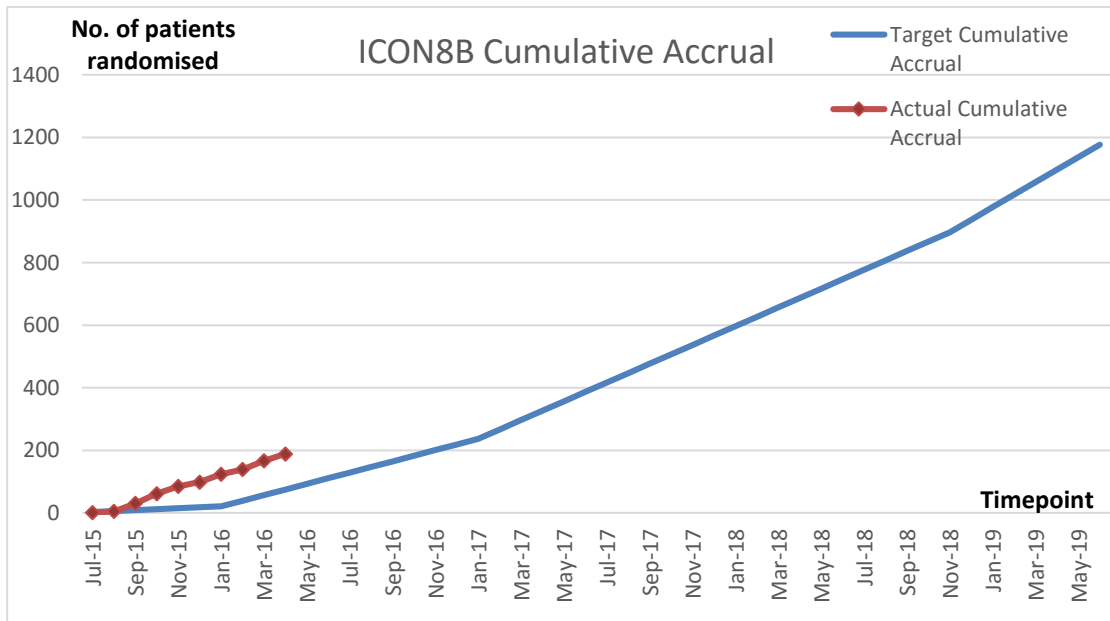
Will be an international trial with participation interest from
Switzerland and Mexico



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ICON8B Trial Progress



First patient recruited
24th July 2015

Accrual and site data up
until 1st May 2016.

Accrual total to date: 196



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ICON8B Expected Vs Actual Monthly Accrual

