**Abstract 971 Figure 2**

**971**

**NK CELL-MEDIATED ERADICATION OF OVARIAN CANCER CELLS WITH A NOVEL CHIMERIC ANTIGEN RECEPTOR DIRECTED AGAINST CD44**

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**Methodology**

We used ovarian cancer cell lines A2780, SKOV3, and OVCAR3 as well as primary ovarian cancer cells (P1, P2 and P3) harvested from ascites samples of an ovarian cancer patient. An anti-CD44 CAR was generated based on a third-generation CAR design containing CD28 and 41BB as codomains. NK-92 cells were equipped with CAR constructs by lentiviral transduction. IFNγ release assays were performed to demonstrate specific activation of engineered NK cells. Live cell impedance analysis with xCELLigence was used to estimate the anti-tumor activity of NK cells. Cytotoxicity assay NK cells and cisplatin were added to A2780 or primary ovarian cancer cells. After treatment, analysis was done by the CellTiter 96® AQueous One Solution Cell Proliferation Assay.

**Result(s)**

NK92 cells equipped with the anti-CD44 CAR (CD44NK) showed specific cytotoxic activity against CD44-positive ovarian cancer cells and primary ovarian cancer cells (figure 1). Specific activation of engineered NK cells was demonstrated by IFNγ secretion assays. Interestingly, CD44NK cells demonstrated cytotoxic activity under cisplatin treatment. The simultaneous treatment with CD44NK and cisplatin showed higher anti-tumor activity compared to a sequential treatment as shown in figure 2.

**Conclusion**

The new anti-CD44 CAR proved specific killing in ovarian cancer cell lines and primary ovarian cancer cells. We showed that CD44NK retained cytotoxicity during cisplatin incubation. The most potent anti-tumor effect was achieved by simultaneous treatment with CD44NK cells and cisplatin. This study will be the basis for further *in vivo* studies and future clinical developments.

**Abstract 977**

**MUCINOUS BORDERLINE OVARIAN TUMORS: PATHOLOGICAL AND PROGNOSTIC STUDY**

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**Methodology**

Mucinous borderline ovarian tumors (MBT) are characterized by an epithelial proliferation similar to those of well-differentiated adenocarcinomas but are distinguished by the absence of stromal invasion. They are often difficult to diagnose histologically. On the one hand, the invasion of the stroma is not always easy to highlight, and on the other hand, the indirect criteria of invasion are not...
unanimously accepted. The work aims to specify the pathological and clinical features and to highlight the prognostic factors of these tumors.

**Methodology** Our study was retrospective and descriptive including 49 cases of primary borderline mucinous tumors of the ovary, diagnosed at the Department of Anatomical Pathology and Cytology of Salah Azaiez Institute, for a period of 27 years, going from 1992 to 2019.

**Result(s)** The mean age of our patients was 48 years old. Histologically, the cases were divided into 34 cases of pure MBT, 13 cases with intraepithelial carcinoma, and 2 cases associating an intraepithelial carcinoma with microinvasion. The majority of our cases were classified FIGO I and only one case was FIGO III. 14 patients received conservative treatment and 32 received radical treatment. The treatment wasn’t specified in 3 patients. The progress was good in the majority of cases. Only one patient had a contralateral recurrence after a follow-up period of 3 years. There was no significant difference regarding the risk of recurrence and risk factors such as age, gestation, hormonal contraception, hormonal status, FIGO stage, presence of peritoneal pseudomyxoma, intraepithelial carcinoma, and microinvasion.

**Conclusion** The prognosis of TMBL depends closely on their FIGO stage, stage I tumors have a good prognosis. The presence of intraepithelial carcinoma does not influence their prognosis. However, it is necessary to multiply samples to avoid missing a carcinomatous focus with an anergic invasion of the stroma which constitutes a poor prognosis factor.

**Introduction/Background** Elevated Alpha-Feto Protein (AFP) in a young female with ovarian mass is virtually diagnostic of Malignant Germ Cell Tumours. We describe a case with outstanding clinical dilemma where the cause of raised AFP remains unsubstantiated.

**Methodology** A 13 year old girl presented with lower abdominal discomfort. Ultrasound examination suggested large left adnexal dermoid cyst. AFP was elevated at 728ng/dL. CT scan showed left adnexal mass and a suspicious small lesion in liver without any other abdominal lesion. She was overweight with grade-2 fatty liver, mildly raised alkaline-phosphatase, hepatomegaly with family history of liver malignancy. A torted left-adnexal smooth mass was removed during surgery. Peritoneal washing, opposite ovary and systematic peritoneal cavity examination were unremarkable. HPE was inconclusive as the tumour was necrotic. After a gap she attended for follow up and on 4th postoperative-month AFP level was 534.84ng/dL.

Further CT and MRI did not reveal any liver lesions. Right ovary had features of poly cystic ovary (PCO). On 5th postoperative-month PET/CT revealed FDG avid 3.5 cm solid-cystic lesion in right adnexa with SUV Max of 5.6, suspicious of malignancy. Patient and family underwent thorough counselling between extent of surgeries vs chemotherapy.

**Result(s)** On second surgical evaluation the right ovary appeared normal and wedge biopsy was benign. Soon after surgery she attained menarche. At 19th month post index surgery, AFP remained elevated; steady at mid-500 ng/dL level without radiological abnormalities. She is on pathway for weight reduction and regular follow up.

**Conclusion** This case report enriches the limited literature on rare reasons for non-hepatic and non-germ cell tumour AFP elevation. Moderate metabolic avidity on PET/CT may signify intense hormonal activities in premenstrual ovary. Causes like Hereditary Persistence of AFP (HPAFP), persistent elevated AFP due to non-hereditary mutations in enhancer and silencer regions of AFP transcription, dietary inflammatory agents and autoimmune neuroinflammation are some of issues which need further research. It is important to recognise these conditions to avoid inappropriate clinical decisions and minimise anxiety level of all concerned. There is need for worldwide registry and in-depth research with genome and exome sequencing to explore raised AFP with unaccommodating classical pathologies.